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(57) Abstract

DNA constructs containing translational control elements are provided. These 5' regulatory segments facilitate high level expression of transgenes introduced into the plastids of higher plants.

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Translation Control Elements for High-level Protein Expression in the Plastids of Higher Plants and Methods of Use Thereof

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This application claims priority from United States Provisional Applications 60/095,163, filed August 3, 1998, 60/112,257, filed December 15, 1998, 60/095,167 filed August 3, 1998, 60/131,611, filed April 29, 1999 and 60/138,764, filed June 11, 1999 under 35 U.S.C.

\$119(e). The entire disclosures of each of the foregoing are incorporated by reference herein.

Pursuant to 35 U.S.C. §202(c) it is acknowledged that the U.S. Government has certain rights in the invention described herein, which was made in part with funds from the National Science Foundation, Grant Number MCB-96-30763.

FIELD OF THE INVENTION

25 This invention relates to the fields of transgenic plants and molecular biology. More specifically, the invention provides vectors targeting the plastid genome which contain translation control elements facilitating high levels of protein expression in the plastids of higher plants. Both monocots and dicots are successfully transformed with the DNA constructs provided herein.

BACKGROUND OF THE INVENTION

Several publications are referenced in this application in order to more fully describe the state of

the art to which this invention pertains. The disclosure of each of these publications is incorporated by reference herein.

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The chloroplasts of higher plants accumulate individual components of the photosynthetic machinery as a relatively large fraction of total cellular protein. The best example is the enzyme ribulose-1,5-bisphosphate carboxylase-oxygenase (Rubisco) involved in CO2 fixation which can make up 65% of the total leaf protein (Ellis, R.J. 1979). Because of the potentially attainable high protein levels, there is significant interest in exploring chloroplasts as an alternative system for protein expression. To date, protein levels expressed from transgenes in chloroplasts are below the levels of highly-expressed chloroplast genes. Highest levels reported thus far in leaves are as follows: 1% of neomycin phophotransferase (Carrer et al., 1993); 2.5% β -glucuronidase (Staub and Maliga, 1993) and 3-5% of Bacillus thuringiensis (Bt) crystal toxins (McBride et al., 1995). An alternative system, based on a 20 · nuclear-encoded, plastid-targeted T7 RNA polymerase may offer higher levels of protein expression (McBride t al., 1994), although this yield may come at a price.

In bacteria, the rate limiting step of protein synthesis is usually the initiation of translation, involving the binding of the initiator tRNA (formyl-methionyl-tRNA_f) and mRNA to the 70S ribosome, recognition of the initiator codon, and the precise phasing of the reading frame of the mRNA. Translation initiation depends on three initiation factors (IF1, IF2, IF3) and requires GTP. The 30S subunit is guided to the initiation codon by RNA-RNA base pairing between the 3' of the 16S rRNA and the mRNA ribosome binding site,

or Shine-Dalgarno (SD) sequence, located about 10 nucleotides upstream of the translation initiation codon (Voorma, 1996). RNA-RNA interaction between the "downstream box" (DB), a 15 nt sequence downstream of the AUG translational initiation codon and complementary sequences in the 16S rRNA 3' sequence or anti-downstream box (ADB; nucleotide positions 1469-1483) may also facilitate loading of the mRNA onto the 30S ribosome subunit (Sprengart et al., 1996). In addition, specific protein-RNA interactions may also facilitate translation initiation (Voorma, 1996).

Key components of the prokaryotic translation machinery have been identified in plastids, including homologues of the bacterial IF1, IF2 and IF3 initiation factors and an S1-like ribosomal protein (Stern et al., 1997). Most plastid mRNAs (92%) contain a ribosome binding site or SD sequence: GGAGG, or its truncated tri- or tetranucleotide variant. This sequence is similar to the bacterial SD consensus 5'-UAAGGAGGUGA-3' (Voorma, 1996). High level expression of foreign genes of interest in the plastids of higher plants is extremely desirable. The present invention provides novel genetic translational control elements for use in plastid transformation vectors. Incorporation of these elements into such vectors results in protein expression levels comparable to those observed for highly expressed chloroplast genes in both monocots and dicots.

SUMMARY OF THE INVENTION

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5' genetic regulatory regions contain promoters
with distinct DNA sequence information which facilitates
recognition by the RNA polymerase and translational
control elements which facilitate translation. Both of

these components act together to drive gene expression.

In accordance with the present invention, chimeric 5' regulatory regions have been constructed which incorporate translation control elements. Incorporation of these chimeric 5' regulatory regions into plastid transforming vectors followed by transformation of target plant cells gives rise to dramatically enhanced levels of protein expression. These chimeric 5' regulatory regions may be used to advantage to express foreign genes of interest in a wide range of plant tissues. It is an object of the present invention to provide DNA constructs and methods for stably transforming plastids of multicellular plants containing such promoters.

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15 In one embodiment of the invention recombinant DNA constructs for expressing at least one heterologous protein in the plastids of higher plants are provided. The constructs comprise a 5' regulatory region which includes a promoter element, a leader sequence and a downstream box element operably linked to a coding 20 region of said at least one heterologous protein. The chimeric regulatory region acts to enhance translational efficiency of an mRNA molecule encoded by said DNA construct. Vectors comprising the DNA constructs are also contemplated in the present invention. Exemplary 25 DNA constructs of the invention include the following chimeric regulatory regions: PrnnLatpB+DBwt, PrrnLatpB-DB, PrrnLatpB+DBm, PrrnLclpP+DBwt, PrrnclpP-DB, PrrnLrbcL+DBwt, PrrnLrbcL-DB, PrrnLrbcL+DBm, 30 PrrnLpsbB+DBwt, PrrnLpsbB-DB, PrrnLpsbA+DBwt, PrrnLpsbA-DB, PrrnLpsbA-DB(+GC), PrrnLT7g10+DB/Ec, PrrnLT7g10+DB/pt, and PrrnLT7g10-DB. Downstream box sequences preferred for use in the constructs of the

invention have the following sequences: 5'TCCAGTCACTAGCCCTGCCTTCGGCA'3 and 5'CCCAGTCATGAATCACA AAGTGGTAA'3.

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The 5' regulatory segments of the invention have been successfully employed to drive the expression of the bar gene from S. hydroscopicus in the plastids of higher plants. Synthetic bar genes have also been generated and expressed using the DNA constructs of the present invention. These constructs have been engineered to maximize transgene containment in plastids by incorporating rare codons into the coding region that are not preferred for protein translation in microorganisms and fungi.

In yet another embodiment of the invention, at least one fusion protein is produced utilizing the DNA constructs of the invention. An exemplary fusion protein has a first and second coding region operably linked to the 5' regulatory regions described herein such that production of said fusion protein is regulated by said 5' regulatory region. In one embodiment the first coding region encodes a selectable marker gene and the second coding region encodes a fluorescent molecule to facilitate visualization of transformed plant cells. Vectors comprising a DNA construct encoding such a fusion protein are also within the scope of the present invention. An exemplary fusion protein consists an aadA coding region operably linked to a green fluorescent protein coding region. These moieties may be linked by peptide linkers such as ELVEGKLELVEGLKVA and ELAVEGKLEVA.

Plasmids for transforming the plastids of higher plants, are also included in the present invention. Exemplary plasmids are selected from the group

consisting of pHK30(B), pHK31(B), pHK60, pHK32(B), pHK33(B), pHK34(A), pHK35(A), pHK64(A), pHK36(A), pHK37(A), pHK38(A), pHK39(A), pHK40(A), pHK41(A), pHK42(A), pHK43(A), pMSK56, pMSK57, pMSK48, pMSK49, pMSK35, pMSK53 and pMSK54.

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Transgenic plants, both monocots and dicots harboring the plasmids set forth above are also contemplated to be within the scope of the invention.

In yet another embodiment of the invention, methods are provided for producing transplastomic monocots. method comprises a) obtaining embryogenic cells; exposing said cells to a heterologous DNA molecule under conditions whereby said DNA enters the plastids of said cells, said heterologous DNA molecule encoding at least one exogenous protein, said at least one exogenous protein encoding a selectable marker; c) applying a selection agent to said cells to facilitate sorting of untransformed plastids from transformed plastids, said cells containing transformed plastids surviving and dividing in the presence of said selection agent; d) transferring said surviving cells to selective media to promote plant regeneration and shoot growth; and e) rooting said shoots, thereby producing transplastomic monocot plants. The heterologous DNA molecule may be introduced into the plant cell via a process selected from the group consisting of biolistic bombardment, Agrobacterium-mediated transformation, microinjection and electroporation. In one embodiment of the above described method, protoplasts are obtained from the embryogenic cells and the heterologous DNA molecule is delivered to said protoplasts by exposure to polyethylene glycol. Suitable selection agents for the practice of the methods of the invention are

streptomycin, and paromomycin. Monocot plants which may be transformed using the methods of the invention include but are not limited to maize, millet, sorghum, sugar cane, rice, wheat, barley, oat, rye, and turf grass.

In a preferred embodiment a method for producing transplastomic rice plants is provided. This method entails the following steps: a) obtaining embryogenic calli; b) inducing proliferation of calli on modified CIM medium; c) obtaining embryogenic cell suspensions of said proliferating calli in liquid AA medium;

 d) bombarding said embryogenic cells with microprojectiles coated with plasmid DNA;

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- e) tranferring said bombarded cells to selective liquid

 AA medium; f) transferring said cells surviving in AA

 medium to selective RRM regeneration medium for a time

 period sufficient for green shoots to appear; and
- g) rooting said shoots in a selective MS salt medium.

Plasmids suitable for transforming rice as set forth above include pMSK35 and pMSK53, pMSK54 and pMSK49. Transplastomic rice plants so produced are also contemplated to be within the scope of the invention.

In yet a final embodiment of the invention methods for containing transgenes in transformed plants are provided. An emplary method includes the following steps: a) determining the codon usage in said plant to be transformed and in microbes found in association with said plant; and b) genetically engineering said transgene sequence via the introduction of rare microbial codons to abrogate expression of said transgene in said plant associated microbe. In an exemplary embodiment of the method described immediately above the transgene is a bar gene and said rare codons

are arginine encoding codons selected from the group consisting of AGA and AGG, and transgene is not expressed in *E.coli*.

The following definitions will facilitate the understanding of the subject matter of the present invention:

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<u>Heteroplastomic</u>: refers to the presence of a mixed population of different plastid genomes within a single plastid or in a population of plastids contained in plant cells or tissues.

Homoplastomic: refers to a pure population of plastid genomes, either within a plastid or within a population contained in plant cells and tissues. Homoplastomic plastids, cells or tissues are genetically stable because they contain only one type of plastid genome. Hence, they remain homoplastomic even after the selection pressure has been removed, and selfed progeny are also homoplastomic. For purposes of the present invention, heteroplastomic populations of genomes that are functionally homoplastomic (i.e., contain only minor populations of wild-type DNA or transformed genomes with sequence variations) may be referred to herein as "functionally homoplastomic" or "substantially homoplastomic." These types of cells or tissues can be readily purified to a homoplastomic state by continued selection.

Plastome: the genome of a plastid.

Transplastome: a transformed plastid genome.

<u>Transformation of plastids</u>: stable integration of transforming DNA into the plastid genome that is transmitted to the seed progeny of plants containing the transformed plastids.

<u>Selectable marker gene</u>: the term "selectable marker gene" refers to a gene that upon expression confers a selective advantage to the plastids and a phenotype by which successfully transformed plastids or cells or tissues carrying the transformed plastid can be identified.

<u>Transforming DNA</u>: refers to homologous DNA, or heterologous DNA flanked by homologous DNA, which when introduced into plastids becomes part of the plastid genome by homologous recombination.

Operably linked: refers to two different regions or two separate genes spliced together in a construct such that both regions will function to promote gene expression and/or protein translation.

The detailed description as follows provides examples of preferred methods for making and using the DNA constructs of the present invention and for practicing the methods of the invention. Any molecular cloning and recombinant DNA techniques not specifically described are carried out by standard methods, as generally set forth, for example in Sambrook et al., "DNA Cloning, A Laboratory Manual," Cold Spring Harbor Laboratory, 1989.

BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1A. Plastid mRNAs and the small (16S) ribosomal RNA contain complementary sequences downstream of AUG implicating interactions between mRNA and 16S rRNA during translation initiation in plastids. Proposed model is based on data in E. coli (Sprengart et al., 1996); for sequence of 16S rRNA see ref. (Shinozaki et al., 1986b). SD, Shine-Dalgarno sequence; ASD, anti SD region; DB, downstream box; ADB, anti DB region. Watson-

Crick (line) and G-U (closed circle) pairing are marked.

Figure 1B. Sequence of the anti-downstream-box regions (ADB sequence underlined) of the 16S rRNA in plastids (pt; this application) and in E. coli (Ec; Sprengart et al., 1996). The E. coli ADB box contains sequences between nucleotides 1469-1483 of the 16S rRNA (Sprengart et al., 1996), corresponding to nucleotides 1416-1430 of the tobacco 16S rRNA (Dams et al., 1988; sequence between nucleotides 104173-104187 in Shinozaki et al., 1986).

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Figure 2A. Base-pairing between plastid ADB and atpB, clpP, rbcL, psbB and psbA mRNAs (underlined). Multiple alternative DB-ADB interactions are shown. Nucleotides changed to reduce or alter mRNA-rRNA interaction are in lower case. The number of potential nucleotide pairs formed with the 26 nt ADB region is in parenthesis. The number of pairing events affected by mutagenesis is in bold.

Figure 2B. Complementarity of Prrn T7 phage gene 10 leader derivatives with the *E. coli* and plastid ADB sequences. Nucleotides changed to reduce or alter mRNA-rRNA interaction are in lower case. The number of potential nucleotide pairs formed with the 26 nt ADB region is in parenthesis.

Figure 3A. DNA sequence of the chimeric Prrn plastid promoter fragments with atpB and clpP translation control regions. The plasmid name that is the source of the promoter fragment is given in parenthesis. The Prrn promoter sequence is underlined; nucleotide at which transcription initiates in tobacco plastids is marked with filled circle; translational

initiation codon (ATG) is in bold; SD is underlined with a wavy line; nucleotides of the 5' and 3' restriction sites and point mutations are in lower case.

Figure 3B. DNA sequence of the chimeric Prrn plastid promoter fragments with rbcL and psbB translation control regions. For details see description of Fig. 3A.

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Figure 3C. DNA sequence of the chimeric Prrn plastid promoter fragments with psbA translation control regions. For details see description of Fig. 3A.

Figure 3D. DNA sequence of the chimeric Prrn plastid promoter fragments with the T7 phage gene 10 (PrrnLT7g10+DB/Ec) plastid (PrrnLT7g10+DB/pt) and synthetic DB (PrrnLT7g10-DB). For details see description of Fig. 3A.

Figure 4A. Plastid transformation vector pPRV111A with chimeric neo genes. Plasmid serial numbers, for example pHK34, designate pPRV111A plastid transformation vectors derivatives; adjacent plasmid numbers in parenthesis (e.g. pHK14) designate the source of the chimeric neo gene in pUC118 or pBSIIKS+ vectors. Arrows mark orientation of the selectable marker gene (aadA) and of the chimeric neo gene. Plastid targeting sequences are underlined in bold. Components of the chimeric neo genes are: Prrn, rRNA operon promoter fragment; L, leader sequence; DB, downstream box; NheI site which serves as a synthetic DB is marked by a heavy line; neo, neomycin phosphotransferase coding region; TrbcL, rbcL 3'-untranslated region. 16SrDNA, trnV, rps12/7 are plastid genes (Shinozaki et al., 1986). The restriction sites marked for: EcoRI, SphI, StuI, SacI, NheI, NcoI, XbaI, HindIII, BamHI and BglII. Restriction

sites in brackets were eliminated during construction. The neo translation initiation in plasmid pHK36 is included in NcoI site (not marked). The presence and relative order of NheI (**) and NcoI (*) restriction sites in the plasmid pPRV111A -DB derivatives (pHK35, pHK37, pHK40, pHK42, pHK43) are marked by asterisks. The promoter sequences are shown in Figures 3B, C and D.

Figure 4B. Plastid transformation vector pPRV111B with chimeric neo genes. See description of Fig. 4A. The promoter sequences are shown in Fig. 3A.

Figure 5. Construction of Prrn promoter-plastid leader fragments by overlap extension PCR.

PrrnLT7g10+DB/Ec promoter (SacI-NheI fragment) in plasmid pHK18.

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Figure 7. Construction by PCR of the
20 PrrnLT7g10+DB/pt promoter (SacI-NheI fragment) in
plasmid pHK19.

Figure 8. Restriction map of plasmids pHK2 and pHK3 with the Prrn(L)rbcL(S)::neo::TrbcL gene. Restriction enzyme cleavage sites are marked for: BamHI, EcoRI, HindIII, NcoI, NheI, SacI, XbaI.

Figure 9. DNA sequence of the

Prrn(L)rbcL(S)::neo::TrbcL gene in plasmid pHK3. Plasmid

pHK2 carries an identical neo gene, except that there is

an EcoRI site upstream of the SacI site.

Figure 10. NPTII accumulation in tobacco leaves

detected by protein gel blot analysis. Amount of total soluble leaf protein (μ g) loaded on SDS-PAGE gel is indicated above the lanes. Lanes are designated with plasmid used for plant transformation; μ g protein loaded per lane is given below. NPTII standard and Nt-pTNH32 extracts were run as positive controls; extracts from wild-type non-transformed plants (wt) were used as negative controls.

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Figure 11. The levels of neo mRNA in the transplastomic leaves. The blots were probed for neo (top) and cytoplasmic 25S rRNA as loading control (bottom). Positions of the monocistronic neo mRNA in vector pPRV111A (Figure 4A), the monocistronic neo and dicistronic neo-aadA transcripts in vector pPRV111B (Figure 4B) and the monocistronic neo and dicistronic rbcL-neo transcripts in pTNH32 transformed plants (Carrer et al., 1993) are marked. Lanes are designated with the transgenic plant serial number. 4 μg total cellular RNA was loaded per lane.

Figure 12. Fraction of a codon encoding a particular amino acid and triplet frequency per 1000 codons in the mutagenized atpB and rbcL DB region.

Altered nucleotides are in lower case.

Figure 13A. NPTII accumulation in tobacco roots detected by protein gel blot analysis. Lanes are designated with the plasmid used for plant transformation; μ g protein loaded per lane is given below. NPTII standard was run as positive control; extracts from wild-type non-transformed plants (wt) were used as negative controls.

Figure 13B. Steady-state levels of neo mRNA in tobacco roots. The neo probe detects a monocistronic mRNA in plants transformed with vector pPRV111A (Figure 4A), and a monocistronic neo and a dicistronic neo-aadA transcript in plants transformed with vector pPRV111B (Figure 4B). Lanes are designated with the transgenic plant serial number. 4 μ g total cellular RNA was loaded per lane.

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10 Figure 14. Protein gel blot analysis to detect
NPTII accumulation in tobacco seeds. Lanes are
designated with plasmid used for plant transformation;
μg protein loaded per lane is given below. NPTII
standard was run as positive control; extracts from
wild-type non-transformed plants (wt) were used as
negative controls.

Figure 15A. Diagram showing integration of the chimeric neo and aadA genes into the plastid genome by two homologous recombination events via the plastid targeting sequences (underlined). On top is shown a diagram of plasmids pHK30 and pHK32 are plastid transformation vector pPRV111B derivatives (Zoubenko et al., 1994). Horizontal arrows mark gene orientation. For description of chimeric neo genes, see Figure 4B. 16SrDNA, trnV, rps12/7 are plastid genes (Shinozaki et al., 1986). The restriction sites marked for: EcoRI (E), SacI (S), NheI (N), XbaI (X), HindIII (H), BamHI (Ba) and BqlII Restriction sites in brackets were eliminated during construction. In the middle the wild-type plastid DNA region (Wt-ptDNA) targeted for insertion is shown. Lines connecting plasmids and ptDNA mark sites of homologous recombination at the end of the vector

plastid-targeting regions. The transformed plastid genome segment (T-ptDNA) map is shown on the bottom.

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Figure 15B. DNA gel blot analysis confirms of integration of the neo and aadA genes into the plastid genome. The blot on top was probed with the plastid targeting sequence (Probe 1 in Figure 15A). It lights up 4.2-kb and 1.4-kb fragments in transplastomic lines, and a 3.1-kb fragment in wild-type (see Figure 15A). Note that the 1.4-kb signal is week in most clones. The blot on the bottom was probed for neo sequences, which are present only in the transplastomic lines.

Figure 16A. Diagram showing integration of the bar gene into the tobacco plastid genome. Map of the plastid targeting region in plasmid pJEK6 is shown on top. The targeted region of the wild-type plastid genome (wt-ptDNA) is shown in the middle. Integrated transgenes in the transplastome (T-ptDNA) are shown at the bottom. Map positions are shown for: the bar gene; aadA, the selectable spectinomycin resistance gene; 16SrDNA and rps12/7, plastid genes (Shinozaki et al., 1986). Arrows indicate direction of transcription. Map position of the probe (2.5 kb) is marked by a heavy line; the wild-type (2.9-kb) and transgenic (3.3-kb, 1.9-kb) fragments generated by SmaI and BgIII digestion are marked by thin lines.

Figure 16B. DNA gel blot confirms integration of bar into the tobacco plastid genome. Data are shown for transplastomic lines Nt-pJEK6-2A through E, Nt-pJEK6-5A through E and Nt-pJEK6-13A and B, and the wild-type parental line. SmaI-BglII digested total cellular DNA was probed with the 2.5-kb ApaI-BamHI plastid targeting sequence marked with heavy line in Figure 16A.

Figure 17. PAT assay confirms bar expression in tobacco plastids. PAT activity was determined by conversion of PPT into acetyl-PPT using radiolabeled ¹⁴C-Acetyl-CoA. Data are shown for transplastomic lines Nt-pJEK6-2D, Nt-pJEK6-5A and Nt-pJEK6-13B, nuclear transformant Nt-pDM307-10 and wild-type (wt).

Figure 18A. Transplastomic tobacco plants are herbicide resistant. Wild-type and pJEK6-transformed plants 13 days after Liberty spraying (5 ml, 2% solution).

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Figure 18B. Maternal inheritance of PPT resistance in the seed progeny. Seeds from reciprocal crosses with Nt-pJEK6-5A plants germinated on 0, 10 and 50 mg/L PPT. wt x pJEK6-5A, transplastomic used as pollen parent; pJEK6-5A x wt , transplastomic line female parent. Resistant seedlings are green on PPT medium, sensitive seedlings are bleached.

Figure 19. The engineered bacterial bar coding region DNA sequence in plasmid pJEK3 and pJEK6 and encoded amino acid sequence. Nucleotides encoding the rbcL five N-terminal amino acids are in lower case. Nucleotides added at the 3' end during construction are also in lower case. NcoI, BglII and XbaI cloning sites are marked.

Figure 20A. The synthetic bar gene DNA sequence and the encoded amino acid sequence. The arginines encoded by AGA/AGG codons are in bold. Original nucleotides are in capital letters, altered bases are in lower case. Restriction sites used for cloning are marked.

Figure 20B. The synthetic s2-bar gene DNA sequence and the encoded amino acid sequence. The arginines

encoded by AGA/AGG codons are in bold. Original nucleotides are in capital letters, altered bases are in lower case. Restriction sites used for cloning are marked.

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Figure 21. Synthetic and bacterial bar genes. The bar coding region is expressed in the Prrn/TrbcL cassettes. Note that the Prrn promoters differ with respect to the translational control region.

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Figure 22A. PAT is expressed in *E. coli* from bar, but not from s-bar coding region. PAT activity was determined by conversion of PPT into acetyl-PPT using radiolabeled ¹⁴C-Acetyl-CoA. Data are shown for *E. coli* transformed with plasmids pJEK6 and pKO12 carrying the bar gene, and pKO8, carrying s-bar.

Figure 22B. PAT assay confirms expression of bar and s-bar in tobacco plastids. PAT activity was determined by conversion of PPT into acetyl-PPT using radiolabeled ¹⁴C-Acetyl-CoA. Data are shown for transplastomic lines Nt-pJEK6-13B and Nt-pKO3-24a,B carrying bar and s-bar, respectively.

Figure 23A. Plastid transformation vector with FLARE16-S as selectable marker targeting the plastid inverted repeat region. DNA and protein sequence at the aadA-gfp junction. Nucleotides derived from aadA and gfp are in capital, adapters sequences and the point mutation used to create the BstXI restriction site (bold) are in lower case.

Figure 23B. Physical map of plastid transformation vector with FLARE16-S as selectable marker targeting the plastid inverted repeat region. Shown are: the promoter

(P) and 3'UTR (T) of the aadA16pt-gfp coding region and its component parts (aadA and gfp coding regions); rrn16 and rps12/7 plastid genes; restriction endonuclease sites HindIII (removed), SpeI, XbaI, NcoI, BstXI, NheI, EcoRI. In plasmid pMSK56 aadA16pt-gfp is expressed from the Prrn:LatpBDB promoter and encodes FLARE16-S1. In plasmid pMSK57 aadA16pt-gfp is expressed from the Prrn:LrbcLDB promoter and encodes FLARE16-S2.

Figure 24. Localization of FLARE16-S to tobacco plastids by laser scanning confocal microscopy in heteroplastomic tissue. Images were processed to detect FLARE16-S (green) and chlorophyll fluorescence (red) and both in a merged view. Sections are shown from plants expressing FLARE16-S1 (a,b) and FLARE16-S2 (3c-f). Note wild-type and transformed plastids in leaves (3a,c,d), chromoplasts of petals (3b), trichomes (3e) and nongreen root plastids (f). White arrows mark transplastomic organelles. Bars represent 25 μm.

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Figure 25. Immunoblot analysis of FLARE16-S accumulation in chloroplasts. The amount of loaded protein (μg) is indicated above the lanes.

Quantification of FLARE16-S1 (Nt-pMSK56 plants) and FLARE16-S2 (Nt-pMSK57 plants) is based on comparison with a purified GFP dilution series. Extract from a wild-type plant (Nt) was used as negative control.

Figure 26A. Amplification of border fragments confirms integration of FLARE-S genes into the plastid genome. Maps of the plastid targeting regions of the rice (pMSK49) and tobacco (pMSK57) vectors, the segment of the rice and tobacco plastid genomes targeted by the

vectors (Os-wt and Nt-wt), and the same regions after integration of FLARE-S genes. The ends of plastid targeting regions are connected with cognate sequences in the wild-type plastid genome. Plastid genes 16SrDNA, trnV and rps12/7 are marked only in the wild-type plastid genomes. The position of PCR primers (O1-O6) and the PCR fragments generated by them are also shown.

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Figure 26B. Amplification of border fragments confirms integration of FLARE-S genes into the plastid genome. Gels with PCR-amplified left and right border fragments, and with aadA fragment. Results are shown for rice (Os-pMSK49-1 and Os-pMSK49-2) and tobacco (Nt-pMSK57) transplastomic lines and wild-type (Os-wt) rice. The molecular weight markers is EcoRI- and HindIII-digested λ DNA.

Figure 27. Localization of FLARE11-S3 to rice chloroplasts in the Os-pMSK49-5 line by laser scanning confocal microscopy. Images were processed to detect FLARE11-S (green) and chlorophyll fluorescence (red) and both in a merged view. Arrows point to mixed populations of plastids in cells. Bar represents 25 μ m.

- Figure 28. The sequence of FLARE16-S is shown.
- Figure 29. The sequence of FLARE16-S1 is shown.
 - Figure 30. The sequence of FLARE16-S2 is shown.
- 30 Figure 31. The sequence of FLARE11-S is shown.
 - Figure 32. The sequence of FLARE11-S3 is shown.

Figures 33A and 33B. The sequence of pMSK35 is shown.

Figures 34A and 34B. The sequence of pMSK49 is 5 shown.

Figure 35. A table describing the FLARE constructs of the invention.

DETAILED DESCRIPTION OF THE INVENTION

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DNA cassettes for high level protein expression in plastids are provided herein. Higher plant plastid mRNAs contain sequences within 50 nt downstream of AUG that are complementary to the 16S rRNA 3-region. These 15 complementary sequences are approximately at the same position as DB sequences in E. coli mRNAs. See Figures 1A and 2A. Interestingly, the tentative plastid DB sequence significantly deviates from the E. coli DB consensus, since the tobacco plastid and E. coli 16S rRNA sequence in the anti-downstream-box (ADB) region is significantly different (Figure 1B). The feasibility of improving protein expression by incorporating DB sequences in plastids was assessed by constructing a series of chimeric 5' regulatory regions consisting of the plastid rRNA operon o70-type promoter (Prrn-114; Svab and Maliga, 1993; Vera and Sugiura, 1995) and the leader sequence of plastid mRNAs with the native DB, mutagenized DB and synthetic DB sequences. The plastid mRNA leaders differ with respect to the presence and position of the SD sequence. Translation efficiency from the chimeric promoters was determined by expressing the bacterial neo gene in plastids. The neo (or kan) gene encodes neomycin phosphotransferase (NPTII) and

confers resistance to kanamycin in bacteria and plastids (Carrer et al., 1993). We have found that NPTII from the chimeric neo transcripts accumulates in the range of 0.2% to 23% of the total soluble leaf protein, indicating the importance of translational control signals in the mRNA 5' region for high-level protein expression.

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There is great interest in producing recombinant proteins in plants plastids which, thus far have been expressed from nuclear genes only (Arntzen, 1997; Conrad and Fiedler, 1998; Kusnadi et al., 1997). Protein levels produced from the PrrnLrbcL+DBwt and PrrnLT7g10 expression cassettes described here significantly exceed protein levels reported for nuclear genes. Accumulation of NPTII from nuclear genes is typically <<0.1% (Allen et al., 1996), the highest value being 0.4% of the total soluble protein (Houdt et al., 1997). We reported earlier accumulation of 1% NPTII from a plastid neo transgene (Carrer et al., 1993). Other examples for protein accumulation from plastid transgenes are 2.5% β glucuronidase (GUS) (Staub and Maliga, 1993)) and 3-5% of the Bacillus thuringiensis (Bt) crystal toxins (McBride et al., 1995). As compared to this earlier report, we have achieved a significant increase in NPTII levels, up to 23% of total soluble protein.

FLARE-S, a protein obtained by fusing an antibiotic-inactivating enzyme with the Aequorea victoria green fluorescence protein accumulated to 8% and 18% of total soluble protein from the PrrnLatpB+DBwt and PrrnLrbcL+DBwt cassettes provided herein. See Example 8. High-level protein accumulation from the cassettes of the present invention can be clearly attributed to engineering the translational control

region (TCR) of the chimeric genes. These novel genetic elements may be used in different applications to drive expression of proteins with agronomic, industrial or pharmaceutical importance.

5 There is a strong demand for methods that control the flow of transgenes in field crops. Incorporation of the transgenes in the plastid genome rather than the nuclear genome results in natural transgene containment, since plastids are not transmitted via pollen in most 10 crops (Maliga, 1993). Plastid transformation in crops has not been widely employed due to the lack of technology. Enhanced expression of selective markers should yield higher transformation efficiencies. The chimeric promoters of the present invention facilitate 15 extension of plastid transformation to agronomically and industrially important crops. Indeed, high-level expression from the PrrnLatpB+DBwt cassette described here resulted in ~25-fold increase in the frequency of kanamycin-resistant transplastomic tobacco lines. More 20 importantly, high levels of marker gene expression following plastid transformation have been obtained in rice, the first cereal species in which plastid transformation has been successful. The results are set forth in Example 8.

The following examples are provided to illustrate various embodiments of the present invention. They are not intended to limit the invention in any way.

The protocols set forth below are provided to facilitate the practice of the present invention.

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PREPARATION OF CHIMERIC 5' CASSETTES FOR ELEVATED EXPRESSION OF HETEROLOGOUS PROTEINS IN PLASTIDS OF HIGHER PLANTS

Identification of a potential downstream box in plastid mRNAs

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The presence or absence of downstream box elements in mRNA molecules was determined for the following genes: psbB (Tanaka et al., 1987) and psbA (Sugita and Sugiura, 1984), photosystem II genes; rbcL, encoding the 10 large subunit of ribulose-1,5-bisphosphate carboxylase/oxygenase (Shinozaki and Sugiura, 1982); atpB, encoding the ATPase β subunit (Orozco et al., 1990); and clpP, encoding the proteolytic subunit of the Clp ATP-dependent plastid protease (Hajdukiewicz et 15 al., 1997). Interestingly, most or all of the PclpP-53 promoter is downstream of the transcription initiation site, therefore the PrrnLclpP constructs are assumed to contain two promoters: Prrn-114 and PclpP-53. 20 . Transcription initiation sites for these genes were described in references cited above; for nucleotide position of the genes in the plastid genome see

Initially, it was assumed that the plastid ADB is similar in size and position as the *E. coli* ADB in the 16S rRNA. The *E. coli* ADB is localized on a conserved stem structure between nucleotides 1469 to 1483 (15 nt) that corresponds to nucleotides 1416 and 1430 of the plastid 16S rRNA (Dams et al., 1988; Sprengart et al., 1996). Although in both cases, the ADB is contained in the 16S rRNA penultimate stem, the actual ADB sequence is different in plastids and in *E. coli* (Figure 1B). The N-terminal coding regions of plastid genes atpB, clpP, rbcL, petA, psaA, psbA, psbB, psbD and psbE were

Shinozaki et al., 1986.

searched for potential DB sequences. The homology search was carried out with a 26 nucleotide sequence centered on the tentative DB region (Figure 1B). The search revealed short stretches of imperfect homology with alternative solutions. Since the position of DB in the mRNA is quite flexible (Etchegaray and Inouye, 1999), we show four potential DB-ADB interactions for atpB and rbcL in Figure 2A. Two plastid mRNAs were selected to test the role of DB in the translation of plastid mRNAs: 1) atpB mRNA lacks a SD sequence; and 2) rbcL mRNA contains a SD sequence at the prokaryotic consensus. In addition, the phage T7 gene 10 (T7g10) leader was included in the study. This leader has a wellcharacterized E. coli DB sequence (Figure 2B; Sprengart et al., 1996). Additional plastid mRNAs with potential DB sequences shown in Figure 2A are clpP, psbB and psbA.

Experimental strategy to test the efficiency of leader sequences for translation

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To compare the efficiency of translation from the 5'-UTR of the selected genes, the 5'-UTR was cloned downstream of the strong plastid rRNA operon σ^{70} -type promoter (Prrn-114) (Svab and Maliga, 1993; Allison et al., 1996), which initiates transcription from multiple adjacent nucleotides (-114, -113, -111; Sriraman et al., 1998). The promoter fragments were constructed as SacI-NheI or a SacI-NcoI fragments. Construction of the chimeric promoters using conventional molecular biological techniques is set forth in detail in the next section.

Two constructs were prepared for each 5'-UTR selected: one with (+DB) and one without (-DB) a native

downstream box. It will be obvious from the forthcoming discussion, that the -DB constructs have a synthetic DB provided by the NheI restriction site. The promoters were cloned upstream of the coding region of a kanamycin resistance (neo) gene, which is available on an NheI-XbaI or NcoI-XbaI fragment. For the stabilization of the mRNA, the rbcL gene 3'-untranslated region was cloned downstream of neo as an XbaI-HindIII fragment. The chimeric neo genes can therefore be excised from the pUC118 or pBSIIKS+ plasmids as SacI-HindIII fragments. These source plasmids are listed in Table 1.

Table 1. Salient features of chimeric promoters*

15	Source of 5'-UTR (nucleotides from	SD AUG)	DB	Promoter fragment	pUC118(U) or pBSIIKS+(B)	r pPRV111A,B
	atpB (-90/+42)	-	wt	SacI/NheI	pHK10(U)	pHK30 (B)
	atpB (-90/+6)	-	S	SacI/NheI	pHK11(U)	pHK31(B)
20	atpB (-90/42)	-	m	SacI/NheI	pHK50(B)	pHK60(B)
	clpP (-53/+48)	-	wt	SacI/NheI	pHK12(U)	pHK32(B)
	clpP (-53/+6)	-	s	SacI/NheI	рнк13 (Л)	рнк33 (в)
25	rbcL (-58/+42)	+	wt	SacI/NheI	pHK14 (B)	pHK34 (A)
	rbcL (-58/+6)	+	s	SacI/NheI	pHK15 (U).	pHK35 (A)
	rbcL (-58/+42)	+	m	SacI/NheI	pHK54 (B)	pHK64 (A)
	psbB (-54/+45)	+	wt	SacI/NheI ^d	рнк16 (U)	рНК36 (А)
30	psbB (-54/+3)	÷	s	SacI/NcoI ^d	PHK17 (U)	pHK37(A)
	bT7g10+DB/Ec (-63/	+24)+	Ec	SacI/NheI	pHK18 (B)	pHK38(A)
	^b T7g10+DB/pt (-63/	+24)+	pt	SacI/NheI	pHK19(B)	pHK39(A)
35	T7g10-DB (-63/+9)	+	S	SacI/NheI	pHK20 (B)	pHK40(A)
35	psbA (-85/+21)	-	wt	SacI/NheI	pHK21 (U)	pHK41(A)
	psbA (-85/+3)	-	s	SacI/NcoIe	pHK22 (U)	pHK42 (A)
	cpsbA(+GC) (-85/+3)	-	sSacI/NcoIe	_	pHK43 (A)

⁴⁰ a.SD+, SD at prokaryotic consensus position; SD-, no SD at prokaryotic consensus position;

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DB wt, wild-type; m, mutants; s, NheI site as synthetic DB.

bEc or pt refers to construct with E. coli or plastid DB sequence.

spsbA(+GC) indicates addition of GC to the wild-type A at the mRNA 5'-end.

dIn source gene psbB translation initiation codon is within NcoI site; therefor +DB construct pHK16 has this NcoI site upstream of the NheI site; see Figure 9.

^{*}Translation initiation codon is included in NcoI site; NheI site is directly downstream in kan coding region; see Figure 8.

The Prrn promoter fragment is available in plasmid pPRV100A (Zoubenko et al., 1994). The promoters were designed to include sequences between -197 nt and -114 nt upstream of the mature 16S rRNA 5' end. Nucleotide -197 is the 5'-end of the Prrn promoter constructs utilized for these and other studies (Svab and Maliga. 1993; -1 is the first nucleotide upstream of the mature 16S rRNA). The G at the -114 position is one of three transcription initiation sites; the other two are the adjacent C (-113) and A (-111) nucleotides (Allison et al., 1996, Sriraman et al., 1998). The nucleotide at which Prrn transcription would initiate is marked by a filled circle in Figure 3A-D. In most constructs, this is a G (-114) as in the native promoter. In two constructs the G was replaced by an A, as in the psbA promoter which is the source of the leader sequence \$pHK21, pHK22; see below).

20 DESIGN OF THE 5' LEADER FROM atpB

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For the atpB gene, multiple mRNA 5'-ends were mapped in tobacco leaves including at least four primary transcripts indicating transcription from four promoters and a processed 5'-end 90 nucleotides upstream of the translation initiation codon (Orozco et al., 1990). The terminal nucleotide of the processed atpB 5'-end is a G. Therefore, the chimeric PrrnLatpB promoters were designed to initiate transcription at a G, anticipating that the leader sequence of the chimeric transcript will be a perfect reproduction of the processed atpB mRNA 5'-end. Out of the atpB coding region, 42 and 6 nucleotides are included in the +DBwt and -DB constructs, respectively. The 42 nucleotides include

four potential DB sequences shown in Figure 2A. Two point mutations in the leader sequence were designed to eliminate NheI (T to A) and EcoRI (G to A) restriction sites without affecting the predicted mRNA 5' secondary structure. In the -DB constructs, two codons (6 nucleotides) were retained from the native coding region upstream of the NheI restriction site (GCTAGC sequence) in which the stop codon is out-of-frame (Figure 3A). Eleven silent point mutations were introduced in the DB region of the PrrnLatpB+DBm construct to either minimize the number of base pairs, or to change the nature of base pairing (for example G-C to G-U) (Figure 2A; Figure 3A).

DESIGN OF THE 5' LEADER FROM clpP

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Two major mRNA 5'-ends of the clpP gene were mapped in tobacco leaves (Hajdukiewicz et al., 1997). The terminal nucleotide of the proximal primary transcript is a G. Therefore, the chimeric PrrnLclpP promoters were designed to initiate transcription at a G, anticipating that the leader sequence of the chimeric transcript will be a perfect reproduction of the leader transcribed from the Pclp-53 NEP promoter. Out of the clpP coding region, 48 and 6 nucleotides are retained in the +DBwt and -DB constructs, respectively. The 48 nucleotides include four potential DB sequences as shown in Figure 2A. In the -DB constructs, two codons (6 nucleotides) were retained from the native coding region upstream of the NheI restriction site (GCTAGC sequence) in which the stop codon is out-of-frame.

DESIGN OF THE 5' LEADER FROM rbcL

One primary and one processed mRNA 5'-end were mapped in tobacco leaves for the rbcL gene (Shinozaki and Sugiura, 1982). The terminal nucleotide of the 5 processed 5' end is a T. The chimeric PrrnLrbcL promoters were designed to initiate transcription at a G, one nucleotide downstream of the terminal T. Fortytwo and 6 nucleotides out of the rbcL coding region are included in the +DB and -DB constructs, respectively. The 42 nucleotides include four potential DB sequences 10 as shown in Figure 2A. The one point mutation (G to A) in the leader sequence was designed to eliminate an EcoRI restriction site without affecting the predicted mRNA 5' secondary structure. In the -DB constructs, two 15 codons (6 nucleotides) were retained from the native coding region upstream of the NheI restriction site (GCTAGC sequence) in which the stop codon is out-offrame. Twelve silent point mutations were introduced into the DB region of the PrrnLrbcL+DBm construct to either minimize the number of base pairs, or to change 20 the nature of base pairing (for example G-C to G-U) (Figure 2A, Figure 3B).

DESIGN OF THE 5' LEADER FROM psbB

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One primary and one processed mRNA 5'-end for the psbB gene were tentatively identified in tobacco leaves (Tanaka et al., 1987). The leader sequence was designed to initiate transcription from the G (-114) of the Prrn promoter, and include the intact secondary (stem) structure assumed to be involved in stabilizing the mRNA. Forty-five and 3 nucleotides out of the psbB coding region are included in the +DB and -DB constructs, respectively. The 45 nucleotides include

four potential DB sequences shown in Figure 2A. Since the ATG is naturally included in an NcoI site that is used to fuse the neo coding region with the psbB leader, no amino acid from the psbB coding region is added in the -DB construct.

DESIGN OF THE 5' LEADER FROM psbA

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One mRNA 5'-end was mapped for the psbA gene in tobacco leaves (Sugita and Sugiura, 1984). The terminal nucleotide of the primary transcript is an A. Therefore, the chimeric PrrnLpsbA promoters were designed to initiate transcription at an A, anticipating that the leader sequence of the chimeric transcript will be a perfect reproduction of the leader transcribed from the psbA promoter. Twenty-one and 3 nucleotides out of the psbA coding region are included in the +DB and -DB constructs, respectively. The 21 nucleotides include the potential DB sequence as shown in Figure 2A. Since the neo coding region was linked to the chimeric promoter via an NcoI site which includes the translation initiation codon (ATG), no amino acid from the psbA coding region is added in the -DB constructs. This is true of a second -DB promoter, in plasmid PHK23, in which transcription is designed to initiate from the Prrn G (-114) and C (-113) (Figure 3C).

DESIGN OF THE T7 PHAGE GENE 10 LEADER

The T7 phage gene 10 leader (63 nucleotides) was shown to promote efficient translation initiation in E. coli (Olins et al., 1988). This leader is used in the E. coli pET expression vectors (Studier et al., 1990; Novagen Inc.). The terminal nucleotide at the 5'-end is a G. Therefore, the chimeric PrrnT7g10L promoters were

designed to initiate transcription at a G, anticipating that the leader sequence of the chimeric transcript will be a reproduction of the T7 phage gene 10 mRNA, with the exception of a T to A mutation which was introduced to eliminate an XbaI site. Twenty-four and 9 nucleotides from the T7 phage gene 10 coding region are included in the +DB/Ec (with E. coli DB sequence) and -DB constructs, respectively. To compare the efficiency of E. coli and plastid DB sequences in plastids, a second +DB promoter was constructed with the tobacco DB sequence (PrrnT7g10L+DB/pt). The native T7g10 leader has an NheI site directly downstream of the translation initiation codon. This NheI site was removed by a T to A point mutation in the +DB constructs (Figure 3D).

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For introduction into the plastid genome, the chimeric neo genes were cloned into plastid transformation vector pPRV111A or pPRV111B. See U.S. Patent 5,877,402, the disclosure of which is incorporated herein by reference. The pPRV111 vectors 20 target insertions into the inverted repeat region of the tobacco plastid genome, and carry a selectable spectinomcyin (aadA) resistance gene. The sequences of the vectors have been deposited in GenBank (U12812, U12813). The chimeric neo gene in vector pPRV111B is in 25 tandem with the aadA gene, whereas in vector pPRV111A the chimeric neo is oriented divergently. The general outline of the plastid transformation vector with the chimeric neo genes is shown in Figures 4A and 4B.

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CONSTRUCTION OF CHIMERIC Prnn PROMOTERS WITH PLASTID MRNA LEADERS

The chimeric Prrn promoter/leader fragments were constructed as a SacI-NheI or SacI-NcoI fragments (Table 5 1, below) by overlap extension PCR (SOE-PCR), essentially as described in Lefebvre et al., (1995). Construction of the Prrn-plastid leader segments is schematically shown in Figure 5. The objective of the PCR-1 step is to 1) amplify the Prrn promoter fragment 10 while 2) adding a SacI site upstream and a seam-less overlap with the specific downstream leader sequence. The reaction contains: 1) a primer (oligonucleotide) to add a SacI site at the 5'-end of the fragment; 2) a suitable template containing the Prrn promoter sequence in plasmid pPRV100A (Zoubenko et al., 1994); and 3) a 15 primer to add on the overlap with the leader sequence at the 3' of the amplified product. The objective of the PCR-2 step is to create the chimeric promoter with DB sequence using: 1) the product of PCR-1 step as a 20 primer; 2) a suitable DNA template containing the specific leader sequence; and 3) primer (oligonucleotide) to include NheI restriction site at the 3'-end of the amplification product. The product of the PCR-2 is the SacI-NheI chimeric Prrn promoter fragment with DB sequence. The objective of the PCR-3 25 step is to remove the DB sequence while introducing a suitable NheI or NcoI restriction site. The product of PCR-3 is the SacI-NheI or SacI-NcoI chimeric Prrn promoter fragment in which the DB sequence is replaced 30 with the NheI site. The objective of the PCR-4 step is to replace the wild-type DB with a mutant DB. The product of PCR-4 is a SacI-NheI Prrn promoter fragment.

The primers (oligonucleotides) used for the

construction of chimeric promoters are listed in Table 2. The chimeric promoters were obtained by overlap extension PCR using oligonucleotides and DNA templates schematically shown in Figure 5.

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Table 2.

Oligonucleotides used for the construction of chimeric promoters.

#1: 5'-CCCGAGCTCGCTCCCCCGCCGTCGTTC-3'

#2: 5'CGAATTTAAAATAAATGTCCGCTTGCACGTCGATCGGTTAATTCTCCCAGAAATATAGCCATCC-3'

#3: 5'-CCCGCTAGCCGTGGAAACCCCAGAACC-3'

#4: 5'-CCCGCTAGCTCTCATAATAAAAATAAATAAATAATATGTC-3'

#5: 5'-TCACTTTGAGGTGGAAACGTAACTCCCAGAAATATAGCCATCC-3'

#6: 5'-CCCGCTAGCTTCCTCCAGGACTTCG-3'

#7: 5'-CCCGCTAGCAGGCATTAAATGAAAGAAAGAAC-3'

25 #8: 5'-TAAGAATTTTCACAACAACAAGGTCTACTCGACTCCCAGAAATATAGCCATCC-3'

#9: 5'-CCCGCTAGCTTTGAATCCAACACTTGCTTTAG-3'

#10: 5'-CCCGCTAGCTGACATAAATCCCTCCCTAC-3'

#11: 5'-CAAAGATAAATAGACACTACGTAACTTTATTGCATTGCTCCCAGAAATATAGCCATCC-3'

#12: 5'-CCCGCTAGCATCATTCAATACAACGGTATGAACACG-3'

#13: 5'-TTCTAGTGGGAAACCGTTGTGGTCTCCCTCCCAGAAATATAGCCATCC-3'

#14: 5'-CCCGCTAGCCATATGTATATCTCCTTCTTAAAG-3'

40 #15: 5'-CCCGCTAGCCTGTCCACCAGTCATGCTTGCCATA-3'

#16: 5'-CCCGCTAGCCAAGGCAGGGCTAGTGATTGCCATATGTATATCTCCTTC-3'

#17: 5'-TTTGTTTAACTTTAAGAAGGAGATATACATATGGCAAGCATGACTGGTGG-3'

#18: 5'-CTCCTTCTTAAAGTTAAACAAAATTATTTCTAGTGGGAAACCGTTGT-3'

#19: 5'-CAAAATAGAAAATGGAAGGCTTTTTGCTCCCAGAAATATAGCCATCCC-3'

50 #20: 5'-CAAAATAGAAAATGGAAGGCTTTTTTCCCAGAAATATAGCCATCCC-3

#21: 5'-GGGCCATGGTAAAATCTTGGTTTATTTAATC-3'

#22: 5'-GGGGCTAGCTCTCTCTAAAATTGCAGT-3'

#23: 5'-GAATAGCCTCTCCACCCA-3'

5 #24: 5'-CCCGCTAGCCGTGGACACCCCACTTCCACTTGTTGTCGGGTTTATTCTCAT-3'

#25: 5'-CCCGCTAGCTTTGAATCCTACTGAGGCTTTTGTTTCTGTTTGAGGACTCAT-3'

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CONSTRUCTION OF CHIMERIC Prnn PROMOTER/atpB LEADER SEGMENTS

PrrnLatpB+DBwt in plasmid pHK10 (Product of PCR-2) PrrnLatpB-DB in plasmid pHK11 (Product of PCR-3)

PrrnLatpB+DBm in plasmid pHK50 (Product of PCR-4)
PCR-1: Oligonucleotides #1, #2 as primers; plasmid
pPRV100A (Zoubenko et al., 1994) DNA as template.
PCR-2: Product of PCR-1 step, Oligonucleotide #3 as

primers; plasmid pIK79 (see below) DNA as template.

PCR-3: Oligonucleotide #1, #4 as primers; Product of PCR-2 step as template.

PCR-4: Oligonucleotide #1, #24 as primers; Product of PCR-2 step as template.

Plasmid pIK79 is a Bluescript BS+ phagemid derivative which carries a PvuII/XhoI tobacco plastid DNA fragment between nucleotides 55147-60484 containing the rbcL-atpB intergenic region with divergent promoters for these genes (Shinozaki et al., 1986).

30 CONSTRUCTION OF CHIMERIC Prnn PROMOTER/clpP LEADER SEGMENTS

PrrnLclpP+DBwt in plasmid pHK12 (Product of PCR-2)
PrrnLclpP-DB in plasmid pHK13 (Product of PCR-3)
PCR-1: Oligonucleotides #1, #5 as primers; plasmid
pPRV100A (Zoubenko et al., 1994) DNA as template.
PCR-2: Product of PCR-1 step, Oligo #6 as primers;
tobacco Sal8 ptDNA fragment (Shinozaki et al., 1986) as

template.

PCR-3: Oligonucleotide #1, #7 as primers; Product of PCR-2 step as template.

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CONSTRUCTION OF CHIMERIC Prnn PROMOTER/rbcL LEADER SEGMENTS

PrrnLrbcL+DBwt in plasmid pHK14 (Product of PCR-2)
PrrnLrbcL-DB in plasmid pHK15 (Product of PCR-3)

PrrnLrbcL+DBm in plasmid pHK54 (Product of PCR-4)
PCR-1: Oligonucleotides #1, #8 as primers; plasmid
pPRV100A (Zoubenko et al., 1994) DNA as template.
PCR-2: Product of PCR-1 step, Oligonucleotide #9 as primers; plasmid pIK79 DNA (see description of pHK10

PCR-3: Oligonucleotide #1, #10 as primers; Product of PCR-2 step as template.

PCR-4: Oligonucleotide #1, #25 as primers; Product of PCR-2 step as template.

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CONSTRUCTION OF CHIMERIC Prnn PROMOTER/psbB LEADER SEGMENTS

PrrnLpsbB+DBwt in plasmid pHK16 (Product of PCR-2)
PrrnLpsbB-DB in plasmid pHK17 (Promoter from pHK16,

25 digested with SacI/NcoI)

above) as template.

PCR-1: Oligonucleotides #1, #11 as primers; plasmid pPRV100A (Zoubenko et al., 1994) DNA as template.

PCR-2: Product of PCR-1 step, Oligo #12 as primers; tobacco Sal8 ptDNA fragment (Shinozaki et al., 1986) as template.

PCR-3 was not necessary, since the psbB translation initiation codon is naturally included in an NcoI site. Therefore, the -DB derivative could be obtained by

SacI/NcoI digestion of the PCR-2 step.

CONSTRUCTION OF CHIMERIC Prnn PROMOTER/psbA LEADER SEGMENTS

PrrnLpsbA+DBwt in plasmid pHK21 (Product of PCR-2)
PrrnLpsbA -DB in plasmid pHK22 (Product of PCR-3)
PCR-1: Oligonucleotides #1, #20 as primers; plasmid
pPRV100A (Zoubenko et al., 1994) DNA as template.
PCR-2: Product of PCR-1 step, Oligo #22 as primers;
tobacco Sal3 ptDNA fragment (Shinozaki et al., 1986) as template.

PCR-3: Oligonucleotide #1, #21 as primers; Product of PCR-2 step as template.

PrrnLpsbA(GC) -DB in plasmid pHK23 (Product of PCR-2)
PCR-1: Oligonucleotides #1, #19 as primers; plasmid
pPRV100A (Zoubenko et al., 1994) DNA as template.
PCR-2: Product of PCR-1 step, Oligo #21 as primers;
tobacco Sal3 ptDNA fragment (Shinozaki et al., 1986) as
template.

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In all of the above, PCR amplification was carried out with AmpliTaq DNA polymerase (Perkin Elmer) or Pfu DNA polymerase (Stratagene) and "stepdown" PCR that utilizes gradually decreasing annealing temperatures was performed (Hecker and Roux, 1996). The exact amplification conditions for the chimeric Prrn::LatpB promoters are given below. The amplification conditions for the remaining chimeric Prrn - plastid leader promoters were calculated according to Hecker and Roux (1996), and differ only in the annealing temperatures. Description of PCR conditions for the construction of the chimeric Prrn promoters with plastid mRNA leaders is

given below; for interpretation of individual steps see scheme in Figure 5.

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	PCR-1 Program: 50	picomoles of both primer	s per 100 μ l
	1.1 Denature	5 min. at 94 °C	
	2.1 Denature	1 min. at 94 $^{\circ}$ C	
	2.2 Annealing	0.5 min. at 72 °C	3 cycles
10	2.3 Extension	0.5 min. at 72 °C	
	3.1 Denature	1 min. at 94 °C	
	3.2 Annealing	0.5 min. at 69 °C	3 cycles
	3.3 Extension	0.5 min. at 72 °C	
	4.1 Denature	1 min. at 94 $^{\circ}\text{C}$	
15	4.2 Annealing	0.5 min. at 66 °C	3 cycles
	4.3 Extension	0.5 min. at 72 °C	
	5.1 Denature	1 min. at 94 $^{\circ}\text{C}$	•
	5.2 Annealing	0.5 min. at 63 °C	3 cycles
	5.3 Extension	0.5 min. at 72 $^{\circ}\text{C}$	
20	6.1 Denature	1 min. at 94 °C	
	6.2 Annealing	0.5 min. at 60 °C	3 cycles
	6.3 Extension	0.5 min. at 72 °C	·
	7.1 Denature	1 min. at 94 °C	
	7.2 Annealing	0.5 min. at 57 °C	20 cycles
25	7.3 Extension	0.5 min. at 72 $^{\circ}\text{C}$	
	8.1 Extension	10 min. at 72 °C	
	8.2	1 min. at 30 °C	

The PCR-2 program was essentially identical to the PCR1 program set forth above with the following modifications: 1) Primers in 100 μ l were the products of 1st PCR reaction, 50 picomoles of the oligonucleotide primer were used; and 2) the annealing temperature in

stepdown PCR was from 67 °C to 52 °C. Accordingly, the following annealing temperatures were used: Step 2.2, 67 °C; Step 3.2, 64 °C; Step 4.2, 61 °C; Step 5.2, 58 °C; Step 6.2, 55 °C; Step 7.2, 52 °C.

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The PCR-3 and PCR-4 programs were essentially identical to the PCR1 program with the following modification:

1) The annealing temperature in stepdown PCR was from 69 °C to 44 °C. Accordingly, the following annealing temperatures were used: Step 2.2, 69 °C; Step 3.2, 64 °C; Step 4.2, 59 °C; Step 5.2, 54 °C; Step 6.2, 49 °C; Step 7.2, 44 °C. In cases where the yield of the final PCR reaction was too low for efficient cloning, final product was amplified using primers which were used to generate the ends. The final PCR products were digested with the appropriate restriction enzymes (SacI and NheI or SacI and NcoI) and cloned in plasmids pHK2 or pHK3 (see below).

CONSTRUCTION OF CHIMERIC PROMOTERS WITH T7 PHAGE GENE 10 mRNA LEADER SEGMENT

The chimeric Prrn promoter/T7gene10 leader (PrrnLT7g10) fragments were constructed as SacI-NheI fragments (Table 1, below).

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PrrnLT7g10+DB/Ec promoter in plasmid pHK18

In the absence of a proper DNA template, the

PrrnLT7g10+DB/Ec was constructed by employing a modified

polymerase chain reaction (Uchida, 1992) in two PCR

steps, as schematically shown in Figure 6. The PCR-1A

and PCR1B steps generate two fragments in two separate

reactions (A and B). The objective of the PCR-1A step is

to amplify Prrn promoter fragment while: 1) adding a

SacI site upstream (Oligonucleotide #1 in Table 2); and 2) a seam-less overlap with the specific downstream leader sequence (Oligonucleotide #13 in Table 2) using plasmid pPRV100A (Zoubenko et al., 1994) as DNA template. The objective of the PCR-1B step is to amplify part of the T7g10 leader sequence using overlapping oligonucleotides #15 and #17 in Table 2. The NheI site is introduced in oligonucleotide #15. Both PCR-1A and PCR-1B reactions were carried out by stepdown PCR as described above for the construction of the chimeric Prrn promoters.

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PCR-2 reaction generating this chimeric promoter contained:

- a) The products of the PCR-1A and PCR-1B reactions as DNA templates;
 - b) Oligonucleotide #18 (0.5 picomole; Table 2) togenerate overlapping fragments with products of the PCR-1A and PCR-1B reactions;
- 20 c) Oligonucleotides #1 and #15 (Table 2) for amplification of the final product, 50 picomoles each in 100 μ l final volume.

Promoter was amplified by stepdown PCR, as described for the chimeric Prrn promoters above; the annealing temperatures were between 72 °C to 57 °C.

PrrnLT7q10+DB/pt promoter in plasmid pHK19
The promoter fragment was obtained in one PCR step as shown in Figure 7. The reaction contained:

a) The product of the PCR-2 reaction generating promoter PrrnLT7g10+DB/Ec in plasmid pHK18 as DNA template; and b) Oligonucleotides #1 and #16 (Table 2), 50 picomoles each in 100 µl final volume.

Promoter was amplified by stepdown PCR, as described for the construction of chimeric Prrn promoters above; the annealing temperatures were between 72 °C to 52 °C.

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PrrnLT7q10-DB promoter in plasmid pHK20

The promoter fragment was obtained in one PCR step, which is similar to the PCR-3 step in Figure 5. The reaction contained:

a) The product of the PCR-2 reaction generating promoter PrrnLT7g10+DB/Ec in plasmid pHK18 as DNA template; and b) Oligonucleotides #1 and #14 (Table 2), 50 picomoles each in 100 μ l final volume.

Promoter was amplified by stepdown PCR, as described for the chimeric Prrn promoters above; the annealing temperatures were between 72 °C to 52 °C.

The final PCR products were digested with the SacI and NheI restriction enzymes and cloned in plasmid pHK3 to obtain plasmids pHK18, pHK19, pHK20.

Construction of chimeric neo genes

Construction of the chimeric promoters was described in the preceding sections. For determining effects on levels of protein accumulation, the promoters were cloned upstream of a kanamycin-resistance encoding construct, consisting of the neo coding region and the 3'-UTR of the plastid rbcL gene. Such constructs are available in plasmids pHK2 and pHK3, which carry the same Prrn(L)rbcL(S)::neo::TrbcL gene as a SacI-HindIII fragment. Plasmid pHK2 is a pUC118 vector derivative; pHK3 is a pBSIIKS+ derivative. Plasmid maps with relevant restriction sites are shown in Figure 8. DNA

sequence of the neo gene in plasmids pHK2 and pHK3 is shown in Figure 9. Note, that in plasmid pHK2 the neo gene has an EcoRI site upstream of the SacI site (Figure 8). Prrn and TrbcL have been described by Staub and Maliga, 1994; the neo gene derives from plasmid pSC1 (Chaudhuri and Maliga, 1996). The pUC118 and pBSIIKS+ plasmid derivatives which carry the various promoter constructs are listed in Table 1.

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To determine the DNA sequence of the promoter fragments, the plasmids were purified with the QIAGEN Plasmid Purification Kit following the manufacturer's recommendations. DNA sequencing was carried out using a T7 DNA sequencing kit (version 2.0 DNA, Amersham Cat. No. US70770) and primer No. #23 in Table 2, which is complementary to the neo coding sequence. These promoter sequences are shown in Figure 3A-D.

Introduction of chimeric neo genes into the tobacco plastid genome

Suitable vectors are available for the introduction 20 of foreign genes into the tobacco plastid genome. Such vectors are pPRV111A and pPRV111B, which carry a selectable spectinomycin-resistance (aadA) gene and target insertions into the repeated region of the plastid genome (Zoubenko et al., 1994). The chimeric neo 25 genes were cloned into one of these plastid transformation vectors (Table 1) and introduced into the tobacco plastid genome by the biolistic process. From the transformed cells plants were regenerated by standard protocols (Svab and Maliga, 1993). A uniform 30 population of transformed plastid genome copies was confirmed by Southern analysis.

For Southern analysis, total cellular DNA was

prepared by the CTAB method (Saghai-Maroof et al., 1984). Two leaves of each transformed plant were homogenized and incubated at 60°C for 30 minutes in a buffer containing 2% CTAB (tetradecyl-trimethyl-ammonium bromide), 1.4 M NaCl, 20 mM EDTA (pH 8.0), 1 mM Tris/HCl (pH 8.0) and 100 mM β-mercaptoethanol. After chloroform extraction, the DNA was precipitated with isopropyl alcohol and dissolved in water or in TE buffer (10 mM Tris, 1 mM EDTA, pH 8.0). DNA digested with an appropriate restriction enzyme was electrophoresed on 0.8% agarose gel and transferred to nylon membrane using PosiBlot Transfer apparatus (Stratagene). The blots were probed using Rapid Hybridization Buffer and plastid targeting sequences as a probe labeled with random priming (32 P, Boehringer Mannheim Cat No. 1004760).

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Plastid transformation was achieved with each of the plasmids listed in Table 1. Exceptions were plasmids pHK41 and pHK42. It appears that NPTII expression with the psbA leader derivatives was so high that the plants were not viable. It follows that these same leaders may be used to advantage when fused with weaker promoters.

Transplastomic lines are designated by Nt (N. tabacum, the species), the plasmid name (for example pHK30) and an individual line number and a letter identifying regenerated plants. For example, the Nt-pHK30-1D and Nt-pHK30-1C plants were both obtained by transformation with plasmid pHK30, are derived from the same transformation event and were regenerated from the same culture. Nt-pHK30-2 plants are derived from an independent transformation event. Normally, several transformed lines per construct were obtained. However, data are shown here only for one: Nt-pHK30-1D, Nt-pHK31-1C, Nt-pHK60-5A, Nt-pHK32-2F, Nt-pHK33-2A, Nt-pHK34-9C,

Nt-pHK35-4A, Nt-pHK64-3A, Nt-pHK36-1C, Nt-pHK37-2D, Nt-pHK38-2E, Nt-pHK39-3B, Nt-pHK40-12B and Nt-pHK43-1C.

Testing mRNA accumulation by RNA gel blot (Northern) analysis

RNA gel blot analysis was performed to determine steady-state levels of chimeric mRNA in the transplastomic lines. Total leaf RNA was prepared from the leaves and roots of plants grown in sterile culture according to Stiekema et al (1988). RNA (4 μ q per lane) was electrophoresed on 1% agarose gel and transferred to nylon membranes using the PosiBlot Transfer apparatus (Stratagene). The blots were probed using Rapid Hybridization Buffer Amersham) with a 32P-labeled neo probe (Pharmacia, Ready-To-Go Random Priming Kit). The neo probe was obtained by isolating the NheI/XbaI fragment from plasmid pHK2. The template for probing the tobacco cytoplasmic 25S rRNA was a fragment which was PCR amplified from total tobacco cellular DNA with primers 5'-TCACCTGCCGAATCAACTAGC-3' and 5'-GACTTCCCTTGCCTACATTG-3'. RNA hybridization signals were quantified using a Molecular Dynamics PhosphorImager, and normalized to the 25S rRNA signal.

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Testing NPTII accumulation by protein gel blot (Western) analysis

Total soluble protein was extracted from the leaves, roots or seeds of transgenic tobacco plants grown in sterile culture. In case of leaves grown in sterile culture, about 200 mg leaf tissue was homogenized in 1 ml of buffer containing 50 mM Hepes/KOH (pH 7.5), 1 mM EDTA, 10 mM potassium acetate, 5 mM

magnesium acetate, 1 mM dithiothreitol and 2 mM PMSF. The homogenate was centrifuged twice at 4 °C to remove insoluble material. Protein concentration was determined using the Biorad Protein Assay reagent kit. Transgenic tobacco plants expressing neo in the plastid genome (Nt-pTNH32-70, Carrer et al., 1993) and wild type plants were used as positive and negative controls, respectively. Proteins were separated in SDS polyacrylamide gels (SDS-PAGE; 15% acrylamide, 6 M urea) and transferred to nitrocellulose membranes using a semi-dry transfer apparatus (Bio-Rad). After blocking non-specific binding sites, the membrane was incubated with 4,000-fold diluted polyclonal rabbit antiserum raised against NPTII (5Prime-3Prime Inc.). HRPconjugated secondary antibody, diluted 20,000 fold, and ECL chemiluminescence were used for immunoblot detection on X-ray film. NPTII was quantified on the immunoblots by comparison of the experimental samples with a dilution series of commercial NPTII (5Prime-3Prime).

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EXAMPLE 1

DB sequences enhance protein accumulation from rbcL leader; protein accumulation from the atpB translation control signals is high but DB-independent

The role of DB sequences in mRNA translation was tested using neo as the reporter gene. The neo gene encodes the bacterial enzyme neomycin phosphotransferase (NPTII) (Beck et al., 1982). The tested neo genes have the same promoter (Prrn) and transcription terminator (TrbcL), and differ only with respect to the translation control region (TCR) comprising the 5' untranslated region of the mRNA and the coding region N-terminus. Two constructs were prepared with the atpB and rbcL TCRs.

One construct contained the wild-type TCR, including the processed 5' untranslated region and 42 nucleotides of the coding region N-terminus (PrrnLatpB+DBwt, plasmid pHK30, Figure 4B; PrrnLrbcL+DBwt, plasmid pHK34, Figure 5 4A). The second construct contained silent mutations in the 42-nucleotide segment of the atpB and rbcL Nterminal coding regions to either eliminate or alter mRNA and rRNA base pairing (PrrnLatpB+DBm plasmids pHK60, Figure 2A and Figure 4B; PrrnLrbcL+DBm, pHK64, Figure 2A and Figure. 4A). The silent mutations altered 10 the mRNA sequence without effecting the amino acid sequence. For example, 13 potential base pairs may form between the wild-type atpB mRNA and the ADB sequence shown at the bottom in Figure 2A. The 11 silent 15 mutations affect eight base-paring events for this particular ADB-DB interaction. After mutagenesis, there is a possibility for ten base pairing events, most of which are new. The chimeric neo genes were introduced into the tobacco plastid genome by homologous targeting 20 using the biolistic approach (Svab and Maliga, 1993; Zoubenko et al., 1994). NPTII and neo mRNA levels were then assessed in the leaves of transplastomic plants. Since NPTII in wild-type DB-containing and mutant DBcontaining plants has the exact same protein sequence, protein levels in the plants directly reflect the 25 efficiency of mRNA translation. In case of the atpB TCR, mutagenesis of DB reduced protein accumulation to ~4% instead of ~7% (Figure 10 and Table 3). In contrast, mutagenesis of rbcL DB had a dramatic effect, reducing 30 NPTII accumulation 35-fold. Thus, DB-ADB interaction is very important for translation of the plastid rbcL mRNA, but is less important for translation of the atpB mRNA.

We also prepared a third construct set with the atpB and rbcL leaders, but without the native DB (PrrnLatpB-DB, plasmid pHK31, Figure 4B; PrrnLrbcL-DB, plasmid pHK35, Figure 4A). The neo coding region in these constructs is directly linked to the Prrn promoter via a synthetic NheI restriction site. The NheI restriction site (GCTAGC) is fully complementary to the ADB region (Figure 2B), therefore it was hoped that it would function as a DB sequence. Utility of NheI site as an alternative DB could be best judged by NPTII accumulation from the rbcL leader, which is highly dependent on DB. High levels of NPTII from the NheI construct (4.7%) relative to the mutant DB (0.3%) indicate, that linking the coding region via an NheI site provides a suitable DB for expressing foreign polypeptides (Figure 10, Table 3).

TABLE 3

Levels of NPTII and neo mRNA in tobacco leaves

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	SD	DB	NPTII(%)	neo mRNA	NPTII/neo
Nt-pTNH32-70	+	-	2.10±0.33	41.5	5.06
Nt-pHK30-1D	(+)	wt	7.02±0.82	70.05±12.33	8.85
Nt-pHK31-1C	(+)	ន	2.52±0.79	100	2.52
Nt-pHK60-5A	(+)	m	4.03±1.45	91.57±12.76	4.40
Nt-pHK32-2F	-	wt '	1.17 <u>+</u> 0.05	49.33±7.76	2.37
Nt-pHK33-2A	-	s	0.21±0.05	49.55 <u>±</u> 6.67	0.42
Nt-pHK34-9C	+	wt	10.83±3.84	48.91±22.65	22.14
Nt-pHK35-4A	+	s	4.68±1.84	21.41 <u>+</u> 7.88	21.86
Nt-pHK64-3A	+	m	0.31±0.15	52.47±4.29	0.59
Nt-pHK36-1C	+	wt	2.17 <u>±</u> 70.97	68.8	3.15
Nt-pHK37-2D	+	s	2.35±0.05	42.3	5.56

5	Nt-pHK43-1C	(+)	s	0.65±0.28	13.2	4.92
	Nt-pHK40-12B	+	s	23.00±5.40	90.27±31.83	25.48
	Nt-pHK39-3B	+	pt	0.16±0.13	13.12±1.27	1.22
	Nt-phk38-2E	+	EC	16.39±3.42	47.59±19.06	34.44

DISCUSSION

In bacteria, mutagenesis or deletion of the DB 10 reduces translation 2- to 34-fold, depending on the individual mRNA (Etchegaray and Inouye, 1999; Faxen et al., 1991; Ito et al., 1993; Mitta et al., 1997; Sprengart et al., 1996). Furthermore, reliance on the DB increases when the SD sequence is removed (Sprengart et al., 1996; Wu and Janssen, 1996). In our experiments, no 15 variation was made in the atpB or rbcL 5'UTR, only sequences downstream of the AUG were altered. Mutagenesis of the atpB DB region reduced protein levels -2-fold. Although the atpB mRNA does not have a SD 20 directly upstream of AUG, we speculate that it probably has an alternate mechanism for translation initiation that reduces its dependence on the DB. Alternatively translation initiation may be facilitated by activator proteins as described for Chlamydomonas chloroplasts 25 (Rochaix, 1996; Stern et al., 1997). The consequence of DB mutagenesis on rbcL translation was a dramatic 35fold drop in NPTII levels. Accordingly, efficient rbcL translation is highly dependent on DB-ADB interactions. Genes in both prokaryotes and eukaryotes show biases in 30 the usage of the 61 amino acid codons and have a tRNA population closely matched to the overall codon bias of the resident mRNA population. Incorporation of synonymous minor codons in the coding region may dramatically reduce translation (Makrides, 1996) and 35 destabilize the mRNA (Deana et al., 1998). A well-

characterized example for minor codons causing reduced expression in *E. coli* are the AGA/AGG arginine codons recognized by the same tRNA which are present at the frequency of 2.6 and 1.6 per thousand codons.

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Therefore, we have compared codon usage bias and frequency of triplets per 1000 nucleotides in the wildtype and mutagenized atpB and rbcL DB regions. Since we studied NPTII accumulation in leaves, the values shown in Figure 12 were calculated for the highly expressed rbcL, psaA, psaB, psaC, psbA, psbB, psbC, psbD, psbE and psbF photosynthetic genes using the Genetics Computer Group (GCG; Madison Wisconsin) codon frequency program. Codon usage bias and triplet frequency is comparable in the wild-type and mutant DB regions of both atpB and rbcL. In addition, the mRNAs for the wild-type and mutant DB constructs accumulate at similar levels. Therefore, the dramatic change in NPTII acccumulation from the PrrnLrbcL+DBm promoter in the Nt-pHK64 line can not be attributed to incorporation of a rare codon in the mutant DB region.

We have shown here that sequences downstream of the translation initiation codon may dramatically affect mRNA translation. Therefore, silent mutations in the DB region of heterologous proteins may significantly improve expression in chloroplasts by increasing complementarity of the mRNA with the plastid rRNA penultimate stem structure.

There are significant differences in NPTII accumulation from neo transgenes with different leaders and the same synthetic DB (Table 3). This indicates that the 5'UTR is an important determinant of translation efficiency. Many data are available supporting the importance of 5'UTR as a target for translational

control in higher plants (Hirose and Sugiura, 1996; Staub and Maliga, 1993; Staub and Maliga, 1994b) and the unicellular alga *Chlamydomonas* (Mayfield et al., 1994; Nickelsen et al., 1999; Sakamoto et al., 1993; Zerges et al., 1997). The data presented herein demonstrate that translation efficiency in plastids is determined by sequences both upstream and downstream of the AUG.

10 EXAMPLE 2

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Study of phage T7g10 translation control sequences indicates that the efficient DB in plastids has loose complementarity to ADB

Since the actual ADB sequence is different in plastids 15 and E. coli, we anticipated (Sprengart et al., 1996; Etchegaray & Inoyue, 1999) that replacement of the E. coli DB with a perfect plastid DB (100% DB-ADB complementarity) would enhance translation in plastids. We choose the phage T7g10 translational control region 20 for the study since it has a well-characterized E. coli DB. Three Prrn promoter derivatives were constructed. Cassette PrrnLT7g10+DB/Ec consists of Prrn fused with the native T7q10 TCR containing the E. coli DB (plasmid pHK38; Figure 2B, Figure 4A). Cassette PrrnLT7g10+DB/pt 25 consists of the Prrn promoter, T7g10 leader and the perfect tobacco DB (pHK39; Figure 2B, Figure 4A). Cassette PrrnLT7q10-DB has the Prrn promoter and T7g10 leader, but lacks the T7g10 DB sequence (pHK40; Figure 2B, Figure 4A). The neo coding region in these 30 constructs is directly linked to the Prrn promoter via a synthetic NheI restriction site. The neo genes in the three expression cassettes were introduced into tobacco

plastids by transformation (Svab and Maliga, 1993; Zoubenko et al., 1994) and the leaves of transplastomic tobacco were tested for NPTII accumulation and mRNA levels (Figures 10, 11; Table 3).

Surprisingly, NPTII levels from the heterologous T7g10 TCR were higher (Nt-pHK38; ~16%) than the levels obtained from the rbcL TCR (Nt-pHK34; ~11%). We expected that incorporation of the plastid DB with 100% complementarity would further enhance NPTII levels. Instead, we found that plants transformed with the construct having the perfect plastid DB (Nt-pHK39) contained NPTII levels 100-fold lower than the plants expressing NPTII from the E. coli TCR (Nt-pHK38; Figures 10; Table 3). This result suggests that, unlike in E. coli, 100% complementarity reduces, rather than enhances translation efficiency. Indeed, none of the highly expressed plastid genes have a perfect DB sequence (Figure 2A). RNA gel blots shown in Figure 11 indicate that Nt-pHK39 plants with the perfect DB contain ~3-fold less neo mRNA. Therefore, a contributing factor to lower NPTII levels in these plants appears to be a faster mRNA turnover rate. Furthermore, NPTII expressed from the PrrnLT7g10 derivatives differ by the DB-encoded amino acids at the N-terminus. Therefore, differential protein turnover rates may be part of the reason for differences

30 DISCUSSION

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This example utilizing the *rbcL* translation control regions reveals that sequences downstream of the translation initiation codon may dramatically affect

in NPTII accumulation. The highest yield of NPTII (23%)

was obtained with the synthetic, NheI-containing DB

mRNA translation. Therefore, silent mutations in the DB region of heterologous proteins may significantly improve expression in chloroplasts by increasing complementarity of the mRNA with the plastid rRNA penultimate stem structure. However, it appears that perfect complementarity is undesirable, as it may accelerate mRNA turnover and reduce the rate of translation. This finding highlights differences in the translation machinery of plastids and E. coli, in which perfect complementarity enhances translation (Etchegaray and Inouye, 1999; Sprengart et al., 1996). It is possible, however, that shifting the region of complementarity relative to AUG or targeting a slightly different region of the penultimate stem may facilitate highly efficient translation of mRNAs with a perfectly matched DB.

The T7g10 constructs have one or two relatively rare AGC serine codons (4.7 per 1000, Figure 12), one of which is encoded in the NheI site. This codon is present in the Nt-pHK38 and Nt-pHK40 plants, which contain the highest levels of NPTII. Further improvement may be expected by replacing the AGC with an AGT serine codon.

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EXAMPLE 3

The clpP, psbB and psbA TCRs have distinct expression characteristics

NPTII accumulation was studied in transplastomic tobacco carrying the PrrnLclpP promoter derivatives. The PrrnLclpP+DBwt (Nt-pHK32-2F) and PrrnLclpP-DB (Nt-pHK33-2A) plants accumulate 1.2% and 0.2% NPTII in their leaves (Figure 10; Table 3). We have found that over-

expression of clpP 5'-UTR causes a mutant phenotype manifested as pale green leaf color and slower growth. This phenotype is normalized in older plants. We assume that the primary cause of this mutant phenotype is the lack of ClpP protein, the clpP gene product. This mutant phenotype is absent in plants transformed with other 5'UTRs. Therefore we believe, that the mutant phenotype is attributable to competition for a clpP-specific nuclear factor. The clpP gene has two introns.

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Preliminary RNA gel blot analysis reveals reduced levels of mature, monocistronic *clpP* mRNA (~30% of wild-type) and accumulation of intron I-containing *clpP* pre-mRNA in the pale-green leaves. Normalization of phenotype coincides with increase of translatable monocistronic *clpP* mRNA to wild type levels. Over-expression of clpP 5'UTR therefore may interfere with splicing of *clpP* pre-mRNA.

NPTII accumulation was also studied in transplastomic tobacco carrying the PrrnLpsbB promoter derivatives. The PrrnL psbB+DBwt (Nt-pHK36-1C) and PrrnL psbB -DB (Nt-pHK37-2D) plants accumulate 2.2% and 2.4% NPTII in their leaves (Figure 10; Table 3). Thus, the synthetic DB sequence in case of the psbB TCR efficiently replaces the native DB sequence.

Conversely, it may rely on an alternative mechanism for translation initiation.

The Prrn promoter constructs with the pshA leader were obtained as described. However, we have been able to introduce only one of them, PrrnLpshA-DB(+GC) into tobacco plastids in line Nt-pHK43-1C. The Nt-pHK43-1C plants accumulate NPTII at a relatively low level (0.6%; Figure 10, Table 3). It is conceivable that the lack of success in introducing the +DB construct is due to the

dramatically elevated expression level of NPTII which is toxic to the plants.

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DISCUSSION

NPTII levels obtained from PrrnLclpP+DBwt (NtpHK32-2F) promoter are relatively low, only 1.2% of the total soluble protein. However, this promoter is desirable for driving expression of selectable marker genes, as the recovery of transplastomic clones is relatively efficient when the neo gene is expressed from this promoter, as shown in Example 4. Expression of neo from the PrrnLclpP+DBwt promoter does not cause a mutant phenotype in tissue culture. Thus, it is suitable to drive the expression of marker genes, so long as the marker gene is subsequently removed. It appears that competition for a nuclear-encoded factor required for processing the clpP introns gives rise to the reduced expression observed. This intron is absent in the clpP genes in the monocots rice (Hiratsuka et al., 1989) and maize (Maier et al., 1995). The PrrnLclpP+DBwt promoter therefore may be used to advantage in the transformation of monocots. Furthermore, the level of the trans-factor required for clpP intron processing is likely to be expressed at different levels in dicots. We anticipate therefore, that expression of the clpP TCR will have no undesirable consequences in other dicot species. It is also possible that the phenotypic consequences of expressing the clpP TCR in plastids is a property of the tobacco line, N. tabacum cv. Petit Havana utilized herein and is absent in other tobacco lines. This would

make the clpP gene TCR a desirable expression tool in both monocots and dicots.

Both psbB leader derivatives accumulate NPTII at comparable levels (2.2% and 2.4%, respectively; Table 3). This 5' regulatory region is a good alternative to the most commonly used rbcL leader when protein accumulation is required in the ~2% range.

In the past, the psbA promoter and leader construct yielded relatively high levels of expression in leaves (2.5% GUS; Staub and Maliga, 1993). Yet these constructs did not contain psbA DB elements. The present invention describes the generation of chimeric promoters that are suitable to obtain high-level protein expression while elucidating the regulatory role played by DB sequences. Prrn is the strongest known promoter in plastids and consequently provides for high levels of NPTII translation. These elevated levels of NPTII can be toxic to the plant and therefore it is difficult to obtain transplastomic lines with the highest prospective levels of NPTII. An alternative approach involves operably linking the psbA leader to a relatively weak promoter. This approach may generate cassettes which are suitable for obtaining relatively high levels of protein accumulation from relatively low levels of mRNA.

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EXAMPLE 4

NPTII accumulation in roots and seeds

Posttranscriptional regulation is an important mechanism of plastid gene expression (Rochaix, 1996; Stem et al., 1997). Therefore, we expected that NPTII accumulation may be tissue-specific due to regulation of

gene expression at the level of mRNA translation. Thus, NPTII accumulation was tested in roots and seeds.

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Testing of NPTII accumulation in roots was carried out with a subset of transplastomic lines (Table 4). Roots for protein extraction were collected from plants grown in liquid MS salt medium (3% sucrose) in sterile cultures incubated on a shaker to facilitate aeration. Protein was extracted from the roots with the leaf protocol and tested for NPTII accumulation (Figure 13 A). The highest level of NPTII, 0.75%, is found in the roots of plants expressing NPTII from the clpP TCR (PrrnLclpP+DBwt construct; pHK32). The second highest value, 0.3%, was found in the roots of plants transformed with plasmid pHK38 expressing NPTII from the T7q10 TCR (PrnnLT7g10+DB/Ec promoter). The level of NPTII was about the same, approximately 0.1 %, in roots expressing the recombinant protein from the atpB and rbcL TCR in pHK30- and pHK34- transformed plants.

Since plastids in the roots are smaller than in leaves, we expected lower levels of NPTII accumulation in the roots than in the leaves. This was true for all the tested roots, except those of the Nt-pHK32 plants. Interestingly, NPTII from the clpP TCR accumulated at almost the same level in the roots (0.75%, Table 4) as in the leaves (approximately 1%, Table 3). This is likely attributable to high levels of the neo mRNA in the roots (Figure 13B). Since the clpP leader includes the minimal PclpP-53 promoter (Sriraman et al., 1998a; NAR 26: 4874) we speculate, that the relatively high mRNA levels are due to activation of PclpP-53 in roots. High levels of expression make the clpP leader a desirable TCR for protein expression in roots.

The T7g10 leader (pHK38) was the most efficient in roots from which the most NPTII accumulated relative to the mRNA (Table 4). Although in the Nt-pHK38 plants, the neo mRNA was 7-times less than in the Nt-pHK32 plants, NPTII levels were almost as high (approximately 0.30% compared to 0.75%) as in the plastids with the clpP TCR (pHK32). High level NPTII accumulation from the T7g10 TCR in leaves (pHK38, pHK40; Table 3) and in roots (pHK38; Table 4) indicates the general utility of the phage T7g10 translation control region for protein expression in plastids.

Protein accumulation was also studied in seeds harvested from the transgenic plants (Figure 14). Protein levels were 0.05% in plants transformed with pHK32 (clpp TCR), and approximately 0.01% in plants transformed with plasmid pHK30 (atpB TCR). No NPTII was detectable in plants in which neo was introduced in the rbcL TCR-construct (plasmid pHK34), indicating differential protein accumulation which is dependent on the choice of the TCR.

Table 4.

Levels of NPTII and neo mRNA in tobacco roots

Strain	NPTII (%)	neo mRNA (%)	NPTII/neo mRNAx103
Nt-pHK30-1D	0.14±0.05	33.7	4.2
Nt-pHK32-2F	0.75±0.35	100	7.5
Nt-pHK34-9C	0.12±0.03	23.5	5.1
Nt-pHK38-2E	0.31±0.04	13.4	23.1

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EXAMPLE 5

High-level NPTII expression facilitates efficient recovery of transplastomic lines by selection for kanamycin resistance

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The plastid genome of higher plants is a 120-kb to 160-kb double-stranded DNA which is present in 1,900 to 50,000 copies per leaf cell (Bendich, 1987). To obtain genetically stable transplastomic lines every one of the plastid genome copies (ptDNA) should be uniformly altered in a plant. Since integration of foreign DNA always occurs by homologous recombination, plastid transformation vectors contain segments of the plastid genome to target insertions at specific locations. Useful, non-selectable genes are cloned next to the selectable marker genes, which are then introduced into the plastid genome by linkage to the selectable marker gene (Maliga, 1993). Transforming DNA is introduced into plastids by the biolistic process (Svab et al., 1990; Svab and Maliga, 1993) or PEG treatment (Golds et al., 1993; O'Neil et al., 1993). Elimination of wild-type genome copies occurs during repeated cell divisions on a selective medium. The success of transformation depends on the success of selective amplification of the few initially transformed genome copies. Therefore the choice of the antibiotic used for the selective amplification of transformed genome copies and the mechanism by which the plant cells are protected from

antibiotic action is a critical parameter to be considered for successful generation of homoplasmic plants.

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The most commonly used antibiotic for the selection of transplastomic lines is spectinomycin, an inhibitor of protein synthesis on plastid ribosomes. Initially, plastid transformation in tobacco was carried out by selection for resistance based on mutations in the plastid 16S rRNA (Svab et al., 1990). Selection was inefficient, yielding about one transplastomic clone per 50 bombarded samples, probably because the 16S rRNA based mutation in recessive. Recovery of transplastomic lines was enhanced ~100-fold by selection for a dominant marker, spectinomycin resistance based on inactivation by aminoglycoside 3" adenyltransferase encoded in a chimeric aadA gene (Svab and Maliga, 1993). In addition to tobacco, selection for spectinomycin resistance (aadA) could be applied to recover transplastomic lines in Arabidopsis and potato. The aadA gene in plants confers resistance to both spectinomycin and streptomycin. Selection for streptomycin resistance was used for plastid transformation in rice, a species resistant to spectinomycin, after bombardment with a chimeric aadA gene. See Example 8.

The need for an alternative marker gene for plastid manipulation has led to testing kanamycin resistance as a selective marker. A chimeric neo (kan) gene, encoding neomycin phosphotransferase, was suitable to recover transplastomic tobacco lines. However, recovery of transplastomic lines was relatively inefficient, yielding only one transplastomic line in ~25 bombarded leaf samples. Furthermore, for every plastid transformation event ~25 to 50 kanamycin resistant lines

were obtained in which integration of the plastid neo construct into the nuclear genome resulted in kanamycin resistance (Carrer et al., 1993). We report here that the efficiency of recovering transplastomic clones is significantly improved when transforming tobacco chloroplasts with a new neo gene expressed from a promoter with the atpB and clpP translation control region. The number of nuclear transformation events is reduced using the cassettes of the present invention. These improvements make the new neo gene a practical tool for plastid genome manipulations.

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DISCUSSION

The chimeric neo genes described in Examples 1-4 were introduced into plastids by selection for the linked spectinomycin resistance (aadA) gene as their suitability for directly selecting transplastomic lines was unknown. The transplastomic lines listed in Table 3 were then tested for resistance to kanamycin by their ability to proliferate on a medium containing 50 mg/L kanamycin. The RMOP meduim used for testing induces formation of green callus and shoot regeneration in the absence of kanamycin. The tissue culture procedures utilized for this example are described in references Carrer et al., 1993 and Carrer and Maliga, 1995.

On the selctive kanamycin medium only scanty, white callus forms from wild-type leaf section. Formation of green callus and shoots from leaf section of plants transformed with pHK plasmids in Table 3 indicates that accumulation of NPTII confers kanamycin resistance. We set out to test if transplastomic clones can be directly selected by kanamycin resistance after bombardment with

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plasmids pHK30 and pHK32. The results are summarized in Table 5.

Bombardment of 25 tobacco leaves with plasmid pHK30 yielded 45 kanamycin resistant lines on a medium containing 50 mg/L kanamycin. Transplastomic neo lines are expected to be resistant to much higher levels, 500 mg/L of kanamycin (Carrer et al., 1993). In addition, in plasmid pHK30 the neo gene is physically linked to a spectinomycin resistance (aadA) gene. Spectinomycin resistance is manifested as kanamycin resistance: sensitive leaf sections form white callus and no shoots whereas resistant leaf sections form green callus and shoots on a selective medium (500 mg/L) RMOP medium. We assumed therefore, that all transplastomic lines should be resistant to both 500 mg/L of kanamycin and 500 mg/L spectinomycin (Carrer and Maliga, 1995). When applying this test we found that 22 of the 45 lines meet these criteria. Digestion of the plastid DNA with the EcoRI restriction enzyme and probing with the plastid targeting region should detect 3.1-kb fragment in the wild-type and a 4.2-kb and 1.2-kb fragment in transplastomic lines (Figure 15A). DNA gel blot analysis of seven of the kanamycin-spectinomycin resistant lines confirmed integration of both transgenes into the plastid genome (Figure 15B). Therefore, we assume that all 22 kanamycin-spectinomycin lines are transplastomic (Table 5).

Bombardment of 30 tobacco leaves with plasmid pHK32 yielded 28 kanamycin resistant lines on a medium containing 50 mg/L kanamycin. We have identified 11 double-resistant lines by testing these on a medium containing 500 mg/L of kanamycin and 500 mg/L spectinomycin. All six tested were transplastomic by DNA

gel blot analysis (Figure 15B), therefore we believe that all eleven are transplastomic (Table 5).

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TABLE 5

SELECTION OF TRANSPLASTOMIC TOBACCO

10 CLONES BY KANAMYCIN RESISTANCE

	Vector	No.	Kan. Res.	Kan. Res.	Kan. Res.	Transplastomic
		leaves	50 mg/L	500 mg/L	500 mg/L	
15					Spec. Res.	
					500 mg/L	•
	pTNH32	29	59	7		0
		50*	52			2
		25ª	47	4		1
20	pHK30	25	45		22	22
	рНК32	30	28		11	11

(°Carrer et al., 1993)

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DISCUSSION

Plastid transformation efficiency should be comparable, if we target the same region of the plastid genome for insertion, use similar size targeting sequences and the same method of DNA delivery.

Therefore, lower transformation efficiencies obtained by selection for kanamycin resistance with the old chimeric neo genes was likely due to the lack of recovery of translastomic clones by selection. We have found that transformation with neo genes expressed from the

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PrrnLatpB+DBwt and PrrnLclpP+DBwt promoters is as efficient as with the aadA gene. This is a significant technical advance, and will facilitate plastid transformation in crops, in which the regenerable tissues contain non-green plastids. Most important targets are the non-green plastids of cereal crops. Kanamycin selection is widely used to obtain transgenic lines after transformation with chimeric neo genes in dicots. However, kanamycin is an undesirable selective agent in monocots such as cereal tissue cultures. However, NPTII also inactivates paromomycin, which may be used to recover nuclear gene transformants at an extremely high efficiency in cereals. See for example, PCT application WO99/05296.

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EXAMPLE 6

Bacterial bar gene expression in tobacco plastids confers resistance to the herbicide phosphinothricin

Bialaphos, a non-selective herbicide, is a tripeptide composed of two L-alanine residues and an analog of glutamic acid known as phosphinothricin (PPT). While PPT is an inhibitor of glutamine synthetase in both plants and bacteria, the intact tripeptide has little or no inhibitory effect in vitro. Bialaphos is toxic for bacteria and plants, as intracellular peptidases remove the alanine residues and release active PPT. Bialaphos is produced by Streptomyces hygroscopicus. The bacterium is protected from phosphinothricin toxicity by phosphinothricin acetyltransferase (PAT), the bar gene product. This enzyme acetylates phosphinothricin or demethylphosphinothricin (Thompson et al., 1987). PPT resistant crops have been obtained by expressing the S.

hygroscopicus bar gene in the plant nucleus. resistant lines were obtained by direct selection for PPT resistance in culture after Agrobacterium tumefaciens-mediated DNA delivery in tobacco, potato, Brassica napus and Brassica oleracea (De Block et al., 5 1987, 1989). Biolistic DNA delivery of chimeric bar genes has been employed to obtain PPT resistant maize (Spencer et al., 1990), rice (Cao, et al, 1992) and Arabidopsis thaliana (Sawaskaki et al., 1994). Construction of transplastomic tobacco plants, in which 10 PPT resistance is based on the expression of bar from S. hygroscopicus in plastids is described in the present example. The vectors utilized to express the bar gene contain an exemplary chimeric 5' regulatory region as set forth in the previous examples. The following 15 material and methods facilitate the practice of this aspect of the present invention.

Construction of plastid bar gene

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A NcoI/XbaI bar gene fragment was generated by PCR amplification using plasmid of pDM302 (Cao et al., 1992) with the following primers:

P1, 5'-AAACCATGGCACCACAAACAGAGAGCCCAGAACGACGCCC-3';

P2, 5'-AAAATCTAGATCATCAGATCTCGGTGACG-3'.

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The ends of the PCR fragment were blunt ended by treatment with the Klenow Fragment of DNA polymerase I. The fragment was then ligated into the EcoRV site of pBluescript II KS+ (Stratagene, La Jolla, CA) to create plasmid pJEK3. Sequence analysis of pJEK3 plasmid DNA revealed that the XbaI site we intended to create through PCR amplification of pDM302 is absent. See Figure 19. The bar gene has the two translation

termination codons followed by vector sequences. The last 20 bp of pJEK3 are:

CCCGTCACCGAGATCTGATGAtcgaattcctgcagcccgggggatccactagttct aga. The bar sequences are in capital (stop codons underlined), the vector sequences are in lower case (XbaI site underlined). Since there is an XbaI site present in the vector 40 bp from the intended XbaI site, it was not necessary to repair this error. The NcoI-XbaI fragment from plasmid pJEK3 was ligated into NcoI-XbaI digested pGS104 plasmid (Serino and Maliga, 1997) to generate plasmid pJEK6. Plasmid pGS104 carries a Prrn-TrbcL expression cassette in a pPRV111B plastid transformation vector. A map of the plastid targeting region of plasmid pJEK6 is shown in Figure 16A.

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Plastid transformation and plant regeneration

Tobacco (Nicotiana tabacum cv. Petit Havana) plants were grown aseptically on agar-solidified medium containing MS salts (Murashige and Skoog, 1962) and sucrose (30g/1). Leaves were placed abaxial side up on RMOP media for bombardment. The RMOP medium consists of MS salts, N6-benzyladenine (1mg/l), 1-naphthaleneacetic acid (0.1 mg/l), thymine (1mg/l), inositol (100 mg/l), agar (6g/1), pH 5.8, and sucrose (30g/1). The DNA was introduced into chloroplasts on the surface of $1\mu\mathrm{m}$ tungsten particles using the DuPont PDS1000He Biolistic gun (Maliga 1995). Spectinomycin resistant clones were selected on RMOP medium containing 500 μ g/ml spectinomycin dihydrochloride. Resistant shoots were regenerated on the same selective medium and rooted on MS agar medium (Svab and Maliga, 1993). The independently transformed lines are designated by the

transforming plasmid (pJEK6) and a serial number, for example pJEK6-2, pJEK6-5. Plants regenerated from the same transformed line are distinguished by letters, for example pJEK6-2A, pJEK6-2B.

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Southern Blot Analysis

Total cellular DNA was isolated from wild-type and transgenic spectinomycin resistant plants with CTAB (Saghai-Maroof et al., 1984). The DNA was digested with the Sma I and BglII restriction endonucleases, separated on a 0.7% agarose gel and blotted onto a Hybond-N nylon membrane (Amersham, Arlington Heights, IL) by a pressure blotter. The membrane was hybridized overnight with an ApaI/ BamHI fragment labeled with $(\alpha\text{--}^{32}P\text{--})d\text{CTP}$ using a dCTP DNA Labeling Beads Kit (Pharmacia Inc, Piscataway, NJ). The membrane was washed 2 times with 0.1X SSPE, 0.2X SDS at 55°C for 30 minutes. Film was exposed to the membrane for 30 minutes at room temperature.

PAT Assay

The PAT assay was performed as described by Spencer et. al. (1990). Leaf tissue (100 mg) from wild type tobacco (wt), transgenic Nt-pDM307-10 tobacco (a line transformed with the nuclear bar gene in plasmid pDM307; Cao et al., 1992), and plastid bar gene transformants was homogenized in 1 volume of extraction buffer (10 mM Na₂HPO₄, 10 mM NaCl). The supernatant was collected after spinning in a microfuge for 10 minutes. Protein (25 mg) was added to 1 mg/ml PPT and ¹⁴C-labeled Acetyl CoA. The reaction was incubated at 37°C for 30 minutes and the entire reaction was spotted onto a TLC plate. Ascending

chromatography was performed in a 3:2 mixture of 1-propanol and NH₄OH. Film was exposed to the TLC plate overnight at room temperature.

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Herbicide Application

Wild type and transgenic plants were sprayed with 5 ml of a 2% solution of Liberty (AgrEvo, Wilmington, DE) with an aerosol sprayer.

RESULTS AND DISCUSSION

First the bacterial bar gene was converted into a plastid gene by cloning the bar coding region into a plastid expression cassette. This cassette consists of an engineered plastid rRNA operon promoter (Prrn) and TrbcL and the 3' UTR of the plastid rbcL gene for stabilization of the mRNA. The plastid bar gene was then cloned into the plastid transformation vector to yield plasmid pJEK6, and introduced into plastids on the surface of microscopic tungsten particles. The bar gene integrated into the plastid genome by two homologous recombination events via the plastid targeting sequences, as shown in Figure 16A. Selection for the linked aadA (spectinomycin resistance) gene on spectinomycin-containing medium eventually yielded cells which carried a uniformly transformed plastid genome population, which were then regenerated into plants.

Integration of bar and aadA was verified by DNA gel blot analysis. Total cellular DNA of wild-type and transplastomic plants was digested with the SmaI and

BglII restriction enzymes and probed with the 2.9-kb ApaI-BamHI plastid targeting fragment of *N. tabacum* (Figure 16B). The two fragments that were expected for the transgenic plants, 3.3 kb and 1.9 kb, were present in each of the transplastomic samples shown in Figure 16B. Absence of the 2.9 kb wild type fragment indicated, that by the time these plants have been regenerated, the wild-type plastid genome copies have been diluted out on the selective medium.

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To determine if the plastid bar gene has been expressed, leaf extracts were assayed for phosphinothricin acetyltransferase (PAT) activity. Conversion of PPT into acetyl-PPT indicated PAT activity in each of the tested transplastomic lines. Data in Figure 17 are shown for the transplastomic lines Nt-pJEK6-2D, Nt-pJEK6-5A and Nt-pJEK6-13B. Interestingly, PAT activity was significantly (>>10-fold) higher when bar was expressed in the plastids, as compared to the bar gene expressed from the cauliflower mosaic virus 35S promoter in the nucleus of the Nt-pDM307-10 plant.

PAT expression confers resistance to PPT in tissue culture and in the greenhouse. When wild type leaf sections are grown in tissue culture, 10 mg/L PPT completely blocks callus proliferation. This same PPT concentration is suitable for the selection of nuclear transformants after bombardment with the nuclear bar construct in plasmid pDM307. Leaf sections of plants expressing bar in plastids show resistance in the presence of up to 100 mg/L PPT in the culture medium. We have tested PPT resistance in the greenhouse, spraying wild-type and transplastomic plants with Liberty, a commercial formulation of PPT, at the recommended field dose of 2%. As shown in Figure 18A, 13 days after the

treatment, the wild type plants were dead while the transgenic plants thrived. Since then the sprayed plants have flowered and set seed. Figure 18B shows maternal inheritance of PPT resistance. Lack of plastid pollen transmission results in a lack of herbicide resistance in progeny pollinated with transgenic pollen. bacterial bar gene has a high G + C content (68.3%; Genbank Accession No. X17220), while plastid genes have a relatively high A + T content; for example the G + C content of the highly expressed psbA and rbcL genes is 42.7% and 43.7%, respectively (Genbank Accession No. Z00044). Differences in the G + C content are also reflected in the codon usage biases. Interestingly, data presented here indicate that expression of bar from S. hygroscopicus is sufficiently high to confer resistance to field levels of the non-selective herbicide PPT. Furthermore, the PAT enzyme levels obtained in the transplastomic lines are significantly higher than those observed in the nuclear transformant. Therefore, further improvement of the expression levels may be obtained by optimizing the codon usage for plastids as set forth in Example 7.

Advantages of incorporating bar in the plastid genome are containment of herbicide resistance due to the lack of pollen transmission in most crops.

Furthermore, the lack of genetic segregation would simplify back-crossing for the introduction of herbicide resistance into additional breeding lines.

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EXAMPLE 7

A Synthetic bar gene Improves Containment and Enhances Expression in Plastids

The bacterial bar gene was introduced into the tobacco plastid genome by transformation with plasmid pJEK6, as described above in Example 6. In plasmid pJEK6 bar is expressed in a cassette consisting of the Prrn(L)rbcL(S) promoter and TrbcL transcription terminator. This plasmid conferred PPT resistance to plants grown in the presence of PPT in the tissue culture medium, but direct selection for transformed lines was not possible. Although the PAT levels in homoplastomic leaves was high, the amount of PAT produced by the few pJEK6 bar copies during the early stage of plastid transformation was probably insufficient to protect the entire cell.

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To improve bar expression in plastids a synthetic gene was created. The codon usage was modified to mimic that of the average tobacco photosynthetic plastid gene. Changing the codon usage lead to a lowered GC content characteristic of higher plant plastid genes. To assist with cloning, restriction enzyme recognition sequences were removed and added as necessary. Codon usage frequency in bacteria reflects relative tRNA abundance: frequent use of codons for rare tRNAs may significantly reduce translation efficiency. We hoped that differential codon usage in plastids and bacteria would reduce or prevent expression of the synthetic gene in bacteria, thereby reducing the danger of horizontal gene transfer to microorganisms. We also hoped that improved bar expression in our novel promoter cassettes will allow direct selection of plastid transformants on PPTcontaining medium.

Materials and Methods for Example 7

Codon comparisons of photosynthetic (rbcL, psaA, psaB, psaC, psbA, psbB, psbC, psbD, psbE, psbF) plastid genes were compiled using GCG (Genetics Computer Group, Madison, WI). DNA mutations were then introduced into the bacterial bar gene making its codon usage more similar to plastid genes, while removing several restriction enzyme sites that could interfere with cloning. See Figure 28. The synthetic bar gene (s-bar) was obtained by single-step assembly of the entire s-bar gene from 28 oligonucleotides (one 44 nt primer, one 30 nt primer and twenty-six 40 nt primers) using PCR (Stemmer et al., 1995). The top and bottom strands of the primers overlap with each other by 20 nucleotides. NcoI and NheI sites were added at the 5' end and a XbaI site was added at the 3' end through PCR amplification. To obtain the complete s-bar gene, a small aliquot of the assembly PCR product was amplified using primers 1A and 14B. Unchanged nucleotides are in upper case, altered nucleotides are in lower case in the primers listed below.

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Primer 1A CCATGGCTAGCCCAGAAAGAAGACCCGGCCGATATTAGACG
Primer 1B GCATATCAGCTTCTGTAGCACGTCTAATATCGGCCGGTCT
Primer 2A TGCTACAGAAGCTGATATGCCAGCAGTTTGTACAATCGTT
Primer 2B CTTGTTTCTATATAATGGTTAACGATTGTACAAACTGCTG
Primer 3A AACCATTATAATAGAAACAAGTACAGTTAAACTTTAGAACTG
Primer 3B tTCTTGAGGTTCTTGAGGTTCAGAACTTTAGAACTG
Primer 4A AACCTCAAGAACCTCAAGAATGGACTGATGATCCG
Primer 4B AAGGATAGCGCTCTCGTAGACGGACTAGATCATCAGTCCA
Primer 5A TCTACGAGAGCGCTATCCTTGGCTTGTAGCAGAAGTTGAC
Primer 5B GCGATACCAGCTATCCTTGGCTTCTACAAGTTGACC
Primer 6A GGTGAAGTAGCTGACTTCCTTGCATACTCTGGAAGG
Primer 6A CCAATCATATGCATTTCTTCCAAGGGCCCCGCATAT
Primer 7A CAAGAAATGCATATGATTGGACAGCTGAACTGATCAACTGTTA

Primer 7B GtTGaTGaCGtGGtGAaACGTAaACaGTtGAtTCaGCtGT
Primer 8A CGTtTCaCCaCGtCAtCAaCGtACaGGACTtGGtTCtACt
Primer 8B TTCAGtAGaTGtGTaTAtAGaGTaGAaCCaAGtCCtGTaC
Primer 9A CTaTAtACaCAtCTaCTGAAaTCttTGGAGGCACAaGGtT
Primer 9B aACAGCtACaACaCTCTTaAAaCCtTGTGCCTCCAaaGAt
Primer10A TtAAGAGtGTtGTaGCTGTtATaGGatTGCCtAAtGAtCC
Primer10B CtTCaTGCATGCGtACaCtTGGaTCaTTaGGCAatCCtAT
Primer11A aAGtGTaCGCATGCAtGAaGCtCTaGGATATGCtCCaaGa
Primer11B CCtGCaGCCCtCAaCATaCCtCttGGaGCATATCCtAGaG
Primer12A GGtATGtTGaGGGCtGCaGGtTTCAAaCAtGGaAACTGGC
Primer13A ATGAtGTaGGTTTtTGGCAaCTtGATTCCAGTCTTGAAa
Primer13B GtAGaACtGGACGaGGGGTACtGGtAGACTGAATCAAG
Primer14A ACCtCCtCGTCCaGTtCTaCCaGTtACtGAGATCTGATGA
Primer14B tctagaTCATCAGATCTCAGTAACtG

The amplified s-bar coding region was then cloned into a pBSIIKS+ plasmid (Stratagene, La Jolla, CA) and sequenced (Figure 20A). The s-bar gene was cloned into cassettes with the chimeric PrrnLatpB+DBwt,

PrrnLrbcL+DBwt and PrrnLT7g10+DB/Ec promoters. Table 6 sets forth the plasmids used in the practice of this example.

Table6. Plasmids with bar genes.

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Plasmid	Promoter	bar	3'UTR	Vector
pKO5		synthetic		pBSIIKS+
			1	
_		(s-bar)		
pKO3	PrrnLatpB+DBwt	synthetic	TrbcL	pPRV111B
		(s-bar)		
pKO8	PrrnLrbcL+DBwt	synthetic	TrbcL	pPRV111A
		(s-bar)		
pKO17	PrrnLT7g10+DB/	synthetic	TrbcL	pPRV111B
	EC	(s-bar)		
pKO12	PrrnLrbcL+DBwt	bacterial	TrbcL	pPRV111A
		(bar)		

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To provide a suitable cloning site at 3'-end of the bacterial bar gene, the EagI/BglII fragment of s-bar was replaced with the cognate fragment of the bacterial bar coding region. Such a bacterial bar gene is incorporated in plasmid pKO12 (Figure 21). In plasmid pKO12 the first 22 nucleotides of the bacterial bar coding region are replaced with nucleotides from the s-bar.

10 RESULTS

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The engineered bacterial bar gene in pJEK6 is expressed both in E. coli and plants, as shown in the previous example. We were interested to test if modification of the codon affects expression of the sbar gene in plastids and in E. coli. In E. coli, s-bar expression was determined by measuring PAT activity. Extracts were prepared from bacteria carrying plasmids pKO3 and pKO8 expressing s-bar from the PrrnLatpB+DBwt and PrrnLrbcL+DBwt promoters, respectively. The radioactive assay did not detect any activity, although extracts from bacteria transformed with plasmids pJEK6 and pKO12 carrying the bacterial bar genes gave strong signals (Figure 22A). In plasmid pKO12 the first 22 nucleotides of the bacterial bar coding region are replaced with nucleotides from the s-bar. Therefore, lack of expression from the s-bar in E. coli is not due to changes within the first 22 nucleotides.

The s-bar was also introduced into plastids by transformation with vector pKO3. Extracts were prepared from pKO3- and pJEK6-transformed tobacco plants, which carry the s-bar and bar genes, respectively. Extracts from both types of plants contained significant PAT

activity (Figure 22B). Therefore, the synthetic bar is expressed in plastids but not in E. coli.

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Changing the bar gene codon usage abrogated expression of the gene in E. coli. This is likely due to the introduction of the rare AGA and AGG arginine codons in the s-bar coding region. The triplet frequency per thousand nucleotides for AGA and AGG is the lowest in E. coli, reflecting low abundance of the tRNA required for translation of these codons. The minor arginine tRNA Arg (AGG/AGA) has been shown to be a limiting factor in the bacterial expression of several mammalian genes. The coexpression of ArgU (dnaY) gene that encodes for tRNA Arg (AGG/AGA) resulted in high level production of the target protein (Makrides 1996). The bacterial bar gene has 14 arginine codons, none of which are the rare AGA/AGG codons. The s-bar gene has five of them, three of which are located within the first 25 codons. Therefore, the likely explanation for the lack of s-bar expression in E. coli is introduction of the rare AGA and AGG arginine codons in the s-bar coding region.

There are proteins, which are toxic to *E. coli* but their expression is desirable in plastid to which it is not toxic. Engineering of these proteins in *E. coli* poses a problem, since the commonly used PEP plastid promoters are active in *E. coli*, thus the gene will be transcribed and the mRNA translated. Incorporation of minor codons in the coding region will prevent translation of these proteins in *E. coli*. Particularly useful in this regard is conversion of arginine codons to AGA/AGG. If no arginine is present in the N-terminal region, an N-terminal fusion may be designed containing multiple AGA/AGG codons to prevent translation of the mRNA.

Plants under field conditions are associated with microbes living in the soil, on the leaves and inside the plants. Gene flow from plastids to these microorganisms has not been shown. However, it would be an added safety measure to incorporate codons in plastid genes, which are rare in the target microorganisms, but are efficiently translated in plastids. Incorporation of AGA/AGG codons into the selective marker genes and the genes of interest will prevent transfer of genes from plants to microbes, which lack the capacity to efficiently translate the AGA/AGG codons. In case of specific plant-microbe associations, based on differences in codon usage preferences genes could be designed which would be expressed in plastids but not in microbes.

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Attempts to directly select transplastomic clones after bombardment with the s-bar constructs so far has failed. The s-bar coding region in Figure 20A contains frequent and rare codons in proportions characteristic of plastid genes. It is possible, that relatively rare codons in a specific context at a critical stage will prevent recovery of plastid transformation events.

Examples for tissue-specific translation of mRNAs dependent on tRNA availability are known (Zhou et al., 1999). Therefore, we designed a second synthetic bar gene, S2-bar, containing only frequent codons (Figure 20B). Plastid transformation with the s2-bar will enable direct selection of plastid transformation events by PPT resistance.

EXAMPLE 8

FLUORESCENT ANTIBIOTIC RESISTANCE MARKER FOR FACILE IDENTIFICATION OF TRANSPLASTOMIC CLONES IN TOBACCO AND RICE

Plastid transformation in higher plants is accomplished through a gradual process, during which all the 300-10,000 plastid genome copies are uniformly altered. Antibiotic resistance genes incorporated in the plastid genome facilitate maintenance of transplastomes during this process. Given the high number of plastid genome copies in a cell, transformation unavoidably yields chimeric tissues, in which the transplastomic cells need to be identified and regenerated into plants. In chimeric tissue, antibiotic resistance is not cell autonomous: transplastomic and wild-type sectors both are green due to phenotypic masking by the transgenic cells. Novel genes encoding FLARE-S, a fluorescent antibiotic resistance enzyme conferring resistance to spectinomycin and streptomycin, which were obtained by translationally fusing aminoglycoside 3''adenylyltransferase [AAD] with the Aequorea victoria green fluorescent protein (GFP) are provided in the present example. FLARE-S facilitates distinction of transplastomic and wild-type sectors in the chimeric tissue, thereby significantly reducing the time and effort required to obtain genetically stable transplastomic lines. The utility of FLARE-S to select for plastid transformation events was shown by tracking segregation of transplastomic and wild-type plastids in tobacco and rice plants after transformation with FLARE-S plastid vectors and selection for resistance to spectinomycin and streptomycin, respectively.

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Plastid transformation vectors contain a selectable marker gene and passenger gene(s) flanked by homologous plastid targeting sequences (Zoubenko et al., 1994), and are introduced into plastids by biolistic DNA delivery (Svab et al., 1990; Svab and Maliga, 1993) or PEG

treatment (Golds et al., 1993; Koop et al., 1996; O'Neill et al., 1993). The selectable marker genes may encode resistance to spectinomycin, streptomycin or kanamycin. Resistance to the drugs is conferred by the expression of chimeric aadA (Svab and Maliga, 1993) and neo (kan) (Carrer et al., 1993) genes in plastids. These drugs inhibit chlorophyll accumulation and shoot formation on plant regeneration media. The transplastomic lines are identified by the ability to form green shoots on bleached wild-type leaf sections. Obtaining a genetically stable transplastomic line involves cultivation of the cells on a selective medium, during which the cells divide at least 16 to 17 times (Moll et al., 1990). During this time wild type and transformed plastids and plastid genome copies gradually sort out. The extended period of genome and organellar sorting yields chimeric plants consisting of sectors of wild-type and transgenic cells (Maliga, 1993). In the chimeric tissue antibiotic resistance conferred by aadA or neo is not cell autonomous: transplastomic and wildtype sectors are both green due to phenotypic masking by the transgenic tissue. Chimerism necessitates a second cycle of plant regeneration on a selective medium. In the absence of a visual marker this is an inefficient process, involving antibiotic selection and identification of transplastomic plants by PCR or Southern probing. The feasibility of visual identification of transformed sectors greatly reduces the effort required to obtain homoplastomic clones.

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The Aequorea victoria green fluorescent protein (GFP) is a visual marker, allowing direct imaging of the fluorescent gene product in living cells without the need for prolonged and lethal histochemical staining

procedures. Its chromophore forms autocatalytically in the presence of oxygen and fluoresces green when absorbing blue or UV light (Prasher et al., 1992; Chalfie et al., 1994; Heim et al., 1994) (reviewed in ref. Prasher, 1995; Cubitt et al., 1995; Misteli and Spector, 1997). The gfp gene was modified for expression in the plant nucleus by removing a cryptic intron, introducing mutations to enhance brightness and to improve GFP solubility (Pang et al., 1996; Reichel et al., 1996; Rouwendal et al., 1997; Haseloff et al., 1997; Davis and Vierstra, 1998). GFP was used to monitor protein targeting to nucleus, cytoplasm and plastids from nuclear genes (Sheen et al., 1995; Chiu et al., 1996; Kšhler et al., 1997), and to follow virus movement in plants (Baulcombe et al., 1995; Epel et al., 1996). GFP has also been used to detect transient gene expression in plastids (Hibberd et al., 1998).

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The expression of GFP by directly incorporating the gfp gene in the plastid genome is described herein. Incorporation of a visual marker, the GFP protein, in the plastid transformation vectors of the present invention facilitates distinction of spontaneous antibiotic resistant mutants and plastid transformants (Svab et al., 1990). Furthermore, transplastomic sectors in the chimeric tissue can be visually identified, significantly reducing the time and effort required for obtaining genetically stable transplastomic lines. The utility of the GFP marker described here is further enhanced by its fusion with the enzyme aminoglycoside 3''-adenylyltransferase [AAD] conferring spectinomycin and streptomycin resistance to plants. Using a marker gene encoding a bifunctional protein, FLARE-S (fluorescent antibiotic resistance enzyme, spectinomycin

and streptomycin), prevents physical separation of the two genes and simplifies engineering. Furthermore, fluorescent antibiotic resistance genes enables extension of plastid transformation to cereal crops, in which plastid transformation is not associated with a readily identifiable tissue culture phenotype.

The following protocols are provided to facilitate the practice of the present example.

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Construction of tobacco plastid vectors. The

aadA16gfp gene encodes FLARE16-S fusion protein, and

can be excised as an NheI-XbaI fragment from plasmid

pMSK51, a pBSKSII+ derivative (Genbank Accesssion No.

Not yet assigned. The fusion protein was obtained by

cloning gfp (from plasmid pCD3-326F) downstream of aadA

(in plasmid pMSK38), digesting the resulting plasmid

with BstXI (at the 3' end of the aadA coding region) and

NcoI (including the gfp translation initiation codon)

and linking the two coding regions by a BstXI-NcoI

compatible adapter. The adapter was obtained by

annealing oligonucleotides 5'-GTGGGCAAAGAACTTGTTGAA

GGAAAATTGGAGCTAGTAGAAGGTCTTAAAGTCGC-3' and 5'CATGGCGACTTTAAGACCTTCTACTAGCTCCAATTTTCCTTCAACAAGTTCTTTGC
CCACTACC-3'. The adapter connects AAD and GFP with a
peptide of 16 amino acid residues (ELVEGKLELVEGLKVA).

The engineered aadA gene (Chinault et al., 1986) in plasmid pMSK38 (pBSIIKS+ derivative) has NcoI and NheI sites at the 5' end and BstXI and XbaI sites at the 3' end of the gene. The NcoI site includes the translation initiation codon; the NheI and BstXI sites are in the coding region close to the 5' and 3' ends, respectively; the XbaI site is downstream of stop codon. The mutations were introduced by PCR using

oligonucleotides 5'-

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GGCCATGGGGGCTAGCGAAGCGGTGATCGCCGAAGTATCG-3' and 5'-CGAATTCTAGACATTATTTGCCCACTACCTTGGTGATCTC-3'.

The gfp gene in plasmid CD3-326F is the derivative of plasmid psmGFP, encoding the soluble modified version of GFP (accession number U70495) obtained under order number CD3-326 from the Arabidopsis Biological Resource Center, Columbus, OH (Davis and Vierstra, 1998). The gfp gene in plasmid CD3-326F is expressed in the PpsbA /TpsbA expression cassette. The gfp gene in plasmid CD3-326F was obtained through the following steps. The BamHI-SacI fragment from CD3-326 was cloned into pBSKS+ vector to yield plasmid CD3-326A. The SacI site downstream of the coding region was converted into an XbaI site by blunting and linker ligation (5'-GCTCTAGAGC; plasmid CD3-326B). An NcoI site was created to include the translation initiation codon and at the same time the internal NcoI site was removed by PCR amplification of the coding region N-terminus with primers 5'-CCGGATCCAAGGAGATATAACACCATGGCTAGTAAAGGAGAAGAACTTTTC-3 ' and 5'-GTGTTGGCCAAGGAACAGGTAGTTTTCC-3'. The PCRamplified fragment was digested with BamHI and MscI restriction enzymes, and the resulting fragment was used to replace the BamHI-MscI fragment in plasmid CD3-326B to yield plasmid CD3-326C. The gfp coding region was excised from plasmid CD3-326C as an NcoI-XbaI fragment and cloned into a psbA cassette to yield plasmid CD3-326D. PpsbA and TpsbA are the psbA gene promoter and 3'- untranslated region derived from plasmids pJS25 (Staub and Maliga, 1993). TpsbA has been truncated by inserting a HindIII linker downstream of the modified BspHI site (Peter Hajdukiewcz, unpublished). The

PpsbA::gfp::TpsbA gene was excised as an EcoRI-HindIII fragment and cloned into EcoRI and HindIII digested pPRV111A, to yield plasmid CD3-326F.

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The aadAl6gfp coding region from plasmid pMSK51 was introduced into two expression cassettes. In plasmid pMSK53 the aadA16gfp coding region is expressed in the PrrnLrbcL+DBwt/TpsbA cassette, and encodes the FLARE16-S2 protein (fluorescent antibiotic resistance enzyme, spectinomcyin). PrrnLrbcL+DBwt is described in the previous examples and derives from plasmid pHK14. construct contains a chimeric promoter composed of the rrn operon promoter, the rbcL gene leader and downstream box sequence. TpsbA is the psbA gene 3' untranslated region, and functions to stabilize the chimeric mRNA. In plasmid pMSK54 the aadA16gfp coding region is expressed in the PrrnLatpB+DBwt/TpsbA cassette, and encodes the FLARE16-S1 protein. PrrnLatpB+DBwt derives from plasmid pHK10, and is a chimeric promoter composed of the rrn operon promoter, the atpB leader and downstream box sequence. See Examples 1-4.

The chimeric aadA16gfp genes were introduced into the topacco plastid transformation vector pPRV111B (Zoubenko et al., 1994). The aadA gene was excised from plasmid pPRV111B with EcoRI and SpeI restriction enzymes, and replaced with the EcoRI-SpeI fragment from plasmids pMSK53 and pMSK54 to generate plasmids pMSK57 (aadA16gfp-S2) and pMSK56 (aadA16gfp-S1).

Construction of rice plastid vectors. Plasmid pMSK49 is a rice-specific plastid transformation vector which carries the aadA11gfp-S3 gene as the selective marker in the trnV/rps12/7 intergenic region (GenBank Accession Number: Not yet assigned). Plasmid pMSK49

carries the rice SmaI-SnaBI plastid fragment (restriction sites at nucleotides 122488 and 125 878 in the genome Hiratsuka et al., 1989) cloned into a pBSKSII+ (Stratagene) vector after blunting the SacI and KpnI restriction sites. The XbaI site present in the rice plastid DNA fragment (position at nucleotide 125032 in the genome (Hiratsuka et al., 1989) was removed by filling in and religation. Prior to cloning the selective marker the progenitor plasmid was digested with the BglII restriction enzyme giving rise to a deletion of 119 nucleotides between two proximal BglII sites (positions at 124367 and 124491). The aadA11gfp-S3 gene was then cloned in the blunted BglII sites.

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The aadA gene in plasmid pMSK49 was obtained by modifying the aadA gene in plasmid pMSK38 (above) to obtain plasmid pMSK39. The modification involved translationally fusing the aadA gene product at its N-terminus with an epitope of the human c-Myc protein (amino acids 410-419; EQKLISEEDL Kolodziej and Young, 1991). The genetic engineering was performed by ligating an adapter obtained by annealing complementary oligonucleotides with appropriate overhangs into NcoI-NheI digested pMSK38 plasmid. The oligonucleotides were: 5'-CATGGGGGCTAGCGAACAAAACTCATTTCTGAAGAAGACTTGc-3' and 5'-CTAGGCAAGTCTTCTTCAGAAAATGAGTTTTTGTTCGCTAGCCCC-3'.

The aadA11gfp gene encoding FLARE11-S was obtained by linking AAD and GFP with the 11-mer peptide ELAVEGKLEVA. To clone aadA and gfp in the same polycloning site, gfp (EcoRI-HindIII fragment; from plasmid CD3-326F) was cloned downstream of aadA in plasmid pMSK39 to obtain plasmid pMSK41. The two genes were excised together as an NheI-HindIII fragment, and cloned into plasmid pMSK45 to replace a kanamycin-

resistance gene yielding plasmid pMSK48. Plasmid pMSK45 is a derivative of plasmid pMSK35 which carries the PrrnLT7g10+DB/Ec promoter. The promoter consists of the plastid rRNA operon promoter and the leader sequence of the T7 phage gene 10 leader. In plasmid pMSK48, aadA is expressed from the PrrnLT7g10+DB/Ec promoter. The aadA and gfp genes were then translationally fused with an BstXI-NcoI adapter that links the AAD and GFP with an 11-mer peptide. The adapter was obtained by annealing oligonucleotides 5'GTGGGCAAAGAACTTGCAGTTGAAGGAAAATTGGAGGTCGC-3' and 5'CATGGCGACCTCCAATTTTCCTTCAACTGCAAGTTCTTTGCCCACTACC-3', which was ligated into BstXI/NcoI digested pMSK48 plasmid DNA to yield plasmid pMSK49. Plasmid pMSK49 has the rice plastid targeting sequences present in plasmid

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pMSK35.

Tobacco plastid transformation. Tobacco leaves from 4 to 6 weeks old plants were bombarded with DNA-coated tungsten particles using the Dupont PDS1000He Biolistic gun (1100 psi). Transplastomic clones were identified as green shoots regenerating on bleached leaf sections on RMOP medium containing 500mg/L spectinomycin dihydrochloride (Svab abd Maliga, 1993). The spectinomycin resistant shoots were illuminated with UV light (Model B 100AP, UV Products, Upland, California, USA). Shoots emitting green light were transferred to spectinomycin free MS medium (Murashige and Skoog, 1962) (3% sucrose) on which fluorescent (transplastomic) and non- fluorescent (wild-type) sectors formed. Fluorescent sectors were excised, and transferred to selective (500 mg/L spectinomycin) shoot regeneration (RMOP) medium. Regenerated shoots were tested for uniform transformation by Southern analysis.

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Rice plastid transformation. Callus formation from mature Oryza sativa cv. Taipei 309 seeds was induced on a modified CIM medium (Tompson et al., 1986), containing MS salts and vitamins (2 mg/L glycine, 0.5 mg/L nicotinic acid, 0.5 mg/L pyridoxine and 0.1 mg/L thiamine), 2 mg/L 2,4D, 1 mg/L kinetin and 300 mg/L casein enzymatic hydrolysate Type III (Sigma C-1026) and sucrose (30g/L). Embryogenic suspensions from the proliferating embryogenic calli were obtained on the AA medium (Muller and Grafe, 1978). For plastid transformation by the biolistic process rice embryogenic cells were plated on a filter paper on non-selective modified CIM medium (Tompson et al., 1986). The bombarded cells were incubated for 48 hours, transferred to selective liquid AA medium (Muller and Grafe, 1978) (one to two weeks), and then to solid modified RRM regeneration medium (Zhang and Wu, 1988) containing MS salts and vitamins, 100 mg/L myo-inositol, 4 mg/L BAP, 0.5 mg/L IAA, 0.5 mg/L NAA, 30 g/L sucrose and 40 g/L $\,$ maltose and 100 mg/L streptomycin sulfate on which green shoots appeared in two to three weeks. The shoots were rooted on a selective MS salt medium (Murashige and Skoog, 1962) containing 30 g/L sucrose and 100 mg/L streptomycin sulfate. Leaf samples for PCR analysis and confocal microscopy were taken from plants on selective medium.

PCR amplification of border fragments. Total cellular DNA was extracted according to Mettler (Mettler, 1987). The PCR analysis was carried out with a 9:1 mixture of AmpliTaq (Stratagene) and Vent (New England Biolabs) DNA polymerases in the Vent buffer following the manufacturer's recommendations. The left

border fragment was amplified with primers O3 (5'-ATGGATGAACTATACAAATAAG-3'and O4 (5'-GCTCCTATAGTGTGACG-3'). The right border fragment was amplified with primers O5 (5'-ACTACCTCTGATAGTTGAGTCG-3') and O6 (5'-AGAGGTTAATCGTACTCTGG-3'). The aadA part of FLARE-S genes was amplified with primers O1 (5'-GGCTCCGCAGTGGATGGCGGCCTG-3') and O2 (5'-GGCTCCGCAGTGGATGGCGGCCTG-3'). Primer positions are shown in Fig. 5A. Note that the same primers can be used in transplastomic tobacco and rice plants expressing FLARE-S.

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Detection of FLARE-S by fluorescence. FLARE-S expressing sectors in the leaves were visualized by an Olympus SZX stereo microscope equipped for GFP detection with a CCD camera system. Subcellular localization of GFP was verified by laser-scanning confocal microscopy (Sarastro 2000 Confocal Image System, Molecular Dynamics, Sunnyvale, CA). This system includes an argon mixed gas laser with lines at 488 and 568 nm and detector channels. The channels are adjusted for fluorescein and rhodamine images. GFP fluorescence was detected in the FITC channel (488-514 nm). Chlorophyll fluorescence was detected in the TRITC channel (560-580 nm). The images produced by GFP and chlorophyll fluorescence were viewed on a computer screen attached to the microscope and processed using the Adobe PhotoShop software.

Jimmunoblot analysis. Leaves (0.5 g) collected from plants in sterile culture were frozen in liquid nitrogen and ground to a fine powder in a mortar with a pestle.

For protein extraction the powder was transferred to a

centrifuge tube containing 1 ml buffer [50 mM Hepes/KOH (pH 7.5), 1 mM EDTA, 10 mM potassium acetate, 5 mM magnesium acetate, 1 mM dithiothreitol and 2 mM PMSF] and mixed by flicking. The insoluble material was removed by centrifugation at 4°C for 5 min at 11,600 g. Protein concentration in the supernatant was determined using the Biorad protein assay reagent kit. Proteins (20 μl per lane) were separated in 12% SDS-PAGE (Laemmli, 1970). Proteins separated by SDS-PAGE were transferred to a Protran nitrocellulose membrane (Schleicher and Schuell) using a semi-dry electroblotting apparatus (Bio-Rad). The membrane was incubated with Living Colors Peptide Antibody (Clontech) diluted 1 to 200. FLARE-S was visualized using ECL chemilluminescence immunoblot detection on X-ray film. FLARE-S on the blots was quantified by comparison with a dilution series of commercially available purified wild-type GFP (Clontech).

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RESULTS AND DISCUSSION

Tobacco plastid vectors with FLARE-S as the selectable marker.

Two FLARE-S fusion proteins were tested in E. coli. In one, the AAD and GFP were linked by an 11-mer (ELAVEGKLEVA), in the second by a 16-mer (ELVEGKLELVEGLKVA) linker. For transformation in tobacco, the aadA16gfp coding region (16-mer linker) was expressed in two cassettes known to mediate high levels of protein accumulation in plastids. Both utilize the strongest known plastid promoter driving the expression of the ribosomal RNA operon (Prrn), and the 3'-UTR of the highly expressed psbA gene (TpsbA) for the stabilization of the chimeric mRNAs. The PrrnLatpB+wtDB (plasmid pMSK56) and PrrnLrbcL+DBwt (plasmid pMSK57)

promoters utilize the atpB or rbcL gene leader sequences and the coding region N-termini with the downstream box (DB) sequence, respectively. Due to inclusion of the DB sequence in the chimeric genes, the proteins encoded by the two genes are slightly different, having 14 amino acids of the ATP-ase β subunit (atpB gene products) or ribulose 1,5-bisphosphate carboxylase/oxygenase (rbcL gene product) translationally fused with FLARE16-S (FLARE16-S1 and FLARE16-S2, respectively). To obtain a plastid transformation vector with the fluorescent spectinomycin resistance genes, the chimeric genes were cloned into the trnV/rps12/7 plastid intergenic region in plastid vector pPRV111B. Plasmids pMSK56 and pMSK57 (Fig. 23) express FLARE16-S1 and FLARE16-S2, respectively, as markers.

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Identification of transplastomic tobacco clones by fluorescence. Transformation was carried out by biolistic delivery of pMSK56 and pMSK57 plasmid DNA into chloroplast. The bombarded leaves were transferred onto selective (500 mg/L spectinomycin) shoot regeneration medium. Wild-type leaves on this medium bleach and form white callus. Cells with transformed plastids regenerate green shoots. The leaves on the selective medium were regularly inspected with a hand-held long-wave UV lamp for FLARE-S fluorescence.

No fluorescence could be detected in young shoots (3 to 5 mm in size) developing on pMSK56-bombarded leaves. However, formation of bright sectors in the leaves was observed, when these small shoots were transferred onto non-selective plant maintenance medium. In contrast, cultures bombarded with plasmid pMSK57 yielded small fluorescent shoots at an early stage.

These fluorescent shoots, and some of the non-fluorescent ones, developed into plants with bright sectors on non-selective plant maintenance medium. Therefore, FLARE16-S2 is useful for early detection of plastid transformation events. FLARE16-S2 fluorescence in young shoots on a selective medium should be due to relatively high levels of FLARE16-S2. Higher levels of FLARE16-S2 are also indicated by the brighter sectors in variegated leaves expressing FLARE16-S2 as compared to FLARE16-S1.

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The size of sectors was different in individual shoots. FLARE-S expression in different leaf layers was also obvious. With the traditional selection for spectinomycin resistance, the transplastomic and wild-type sectors are not visible. Regeneration of plants with uniformly transformed plastid genomes was greatly facilitated by the fluorescing sectors expressing FLARE-S, which could be readily identified in UV light, dissected, and transferred for a second cycle of plant regeneration on spectinomycin-containing (500 mg/L) selective medium.

Given the high levels of FLARE-S accumulation we were interested to find out, if FLARE-S is toxic to plants. We expected that toxicity should be manifested as lower transformation efficiencies. Bombardment of 30 tobacco leaves with plasmids pMSK56 and pMSK57 yielded 71 and 89 spectinomycin resistant clones, respectively. Out of these, 61 and 77 lines were verified as transplastomic by fluorescence. Plastid transformation in a subset of these was confirmed by confocal laser scanning microscopy (7 clones each; see below) and Southern analysis (4 clones). The frequency of plastid transformation events with the FLARE-S -expressing genes

was slightly higher (~2 instead of ~1 per bombardment) than reported earlier with a chimeric aadA gene at the same insertion site (Svab and Maliga, 1993). Therefore, we assume that accumulation of FLARE-S at high levels is not detrimental. Lack of toxicity is also supported by the apparently normal phenotype of the plants in the greenhouse (not shown).

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Localization of FLARE-S to tobacco plastids by confocal microscopy. Due to phenotypic masking, transplastomic and wild type sectors in a chimeric leaf are both green on a selective medium. However, we have found that in chimeric leaf sectors in the same cell some plastids express FLARE-S while others do not, when observed by confocal microscopy (Fig. 24). FLARE-S and chlorophyll fluorescence were detected separately in the fluorescein and rhodamine channels, respectively. The two images were then overlaid confirming that FLARE-S fluorescence derives from chloroplasts.

Expression of FLARE-S was also studied in non-green plastid types including the chromoplasts in petals and the non-green plastids in root cells (Fig. 24b,f). These studies were carried out in plants, which were homoplastomic for the transgenomes. Homoplastomic state was important, since in non-green tissues chlorophyll could not be used for confirmation of the organelles as plastids. Since FLARE-S expression could be readily detected in chloroplasts as well as non-green plastids, the plastid rRNA operon promoter is apparently active in all plastid types.

FLARE-S accumulation in tobacco leaves.

Accumulation of FLARE-S in homoplastomic leaves was

tested using the commercially available GFP antibody, recognizing the GFP portion (239 amino acid residues) of FLARE16-S (520 amino acids). FLARE16-S1 (532 amino acids) was ~8 %, whereas FLARE16-S2 (532 amino acids) was ~18 % of total soluble leaf protein (Fig. 25). To calculate FLARE16-S concentrations, a GFP dilution series was used as a reference, and the values were than increased by 2.6 to correct for the larger size of the FLARE16-S1 and -S2 proteins.

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expression. In rice, plant regeneration is from nongreen embryogenic cells. Encouraged by FLARE-S
expression in non-green tobacco plastids, we attempted
to transform the non-green plastids of embryogenic rice
tissue-culture cells. Plastid transformation was carried
out using a rice-specific vector expressing FLARE11-S3
and targeting insertion of the aadA11gfp-S3 gene in the
trnV/rps12/7 intergenic region. The location of the
insertion site and the size of plastid targeting
sequences in the rice vector are similar to the tobacco
vectors shown in Fig. 23.

Plastid transformation in rice was carried out by bombardment of embryogenic rice suspension culture cells using gold particles coated with plasmid pMSK49 DNA. Rice cells, as most cereals, are naturally resistant to spectinomycin (Fromm et al., 1987). FLARE-S, however, confers resistance to streptomycin as well (Svab and Maliga, 1993). Therefore, selection for transplastomic lines was carried out on selective streptomycin medium (100 mg/L). Streptomycin at this concentration inhibits the growth of embryogenic rice cells. After bombardment, the rice cells were first

selected in liquid embryogenic AA medium, then on the solid plant regeneration medium, on which the surviving resistant cells regenerated green shoots (12 in 25 bombarded plates). These shoots were rooted, and grown into plants. PCR amplification of border fragments in DNA isolated from the leaves of these plants confirmed integration of aadAllgfp-S3 sequences in the plastid genome (Fig. 26). The left and right border fragments can not be amplified if the gene is integrated into the nuclear genome, as one of the primers (04 or 06) of the pairs is outside the plastid targeting regions.

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FLARE11-S3 expression in the leaves of two of the PCR-positive plants was tested by confocal laser-scanning microscopy. In rice, as in tobacco, the FLARE-S marker confirmed segregation of transplastomic and wild-type plastids (Fig. 27). In rice only a small fraction of chloroplasts expressed FLARE-S. Since individual cells marked with arrows in Fig. 27 contained a mixed population of wild-type and transgenic chloroplasts, FLARE-S in these cells could be expressed only from the plastid genome. Integration of aadA11gfp-S3 into the nuclear genome downstream of plastid-targeting transit peptide would result in uniform expression of FLARE-S in each of the chloroplasts within the cell.

The sequences of the selectable marker genes of the invention are provided in Figures 28-34. Figure 35 depicts a table describing the selectable marker genes disclosed in the present example.

Direct visual identification of transplastomic sectors requires high level expression of FLARE-S in plastids. High GFP expression levels in Arabidopsis were toxic, interfering with plant regeneration. Toxicity of

wild-type (insoluble) GFP was linked to GFP accumulation in the nucleus and cytoplasm, and could be eliminated by targeting it to the endoplasmic reticulum (Haseloff et al., 1997). GFP aggregates were also cytotoxic to E. coli cells (Crameri et al., 1996). To enhance 5 fluorescence intensity and to avoid cytotoxicity, soluble versions of the codon-modified GFP were obtained (Davis and Vierstra, 1998). We have utilized the gene for a soluble-modified GFP described by Davis and Vierstra (Davis and Vierstra, 1998) to create variants 10 of FLARE-S, a fusion protein, which does not have an apparent cytotoxic effect. The frequency of plastid transformation, if affected at all, is increased rather then decreased. In tobacco, we normally obtain one transplastomic clone per bombarded leaf sample (Svab and 15 Maliga, 1993), whereas with the FLARE-S genes on average we could recover two clones per sample. Plant regeneration from highly fluorescent tissue was readily obtained, and the regenerated plants have a phenotype indistinguishable from the wild type. 20

Plastid transformation in rice requires expression of the selective marker in non-green plastids. The rRNA operon has two promoters, one for the eubacterial-type (PEP) and one for the phage-type (NEP) plastid RNA polymerase. The promoter driving FLARE-S expression is recognized only by the eubacterial-type plastid RNA polymerase. Previously, it was assumed that the eubacterial-type promoter is active only in chloroplasts (Maliga, 1998). Accumulation of FLARE-S in roots and petals indicates that PEP is also active in non-green plastids.

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Plastid transformation is a process that unavoidably yields chimeric plants, since cells of

higher plants contain a large number (300 to 50000) of plastid genome copies (Bendich, 1987), out of which initially only a few are transformed. High level expression of FLARE-S in plastids provides the means for 5 visual identification of transplastomic sectors, even if they are present in a chimeric tissue. GFP and AAD could be expressed from two different genes in a plastid transformation vector. However, transformation with a marker gene encoding a bifunctional protein prevents separation of the two genes and simplifies engineering. 10 The fluorescent selective marker will significantly reduce the work required to obtain genetically stable plastid transformants in tcbacco, a species in which plastid transformation is routine. The bottleneck of 15 applying plastid transformation in crop improvement is the lack of technology. In tobacco, chimeric clones with transformed plastids are readily identified by shoot regeneration (Svab et al., 1990). In Arabidopsis, clones with transformed plastids are identified by greening 20 (Sikdar et al., 1998). We have shown here that FLARE-S is a suitable marker to select for transplastomes in embryogenic rice cells, which lack the visually identifiable tissue culture phenotypes exploited in tobacco and Arabidopsis. Data presented here are the 25 first example for stable integration of foreign DNA into the rice plastid genome. These rice plants are heteroplastomic. Uniformly transformed rice plants will be obtained by further selection on streptomycin medium and screening the embryogenic cells for FLARE-S 30 expression. Thus, the FLARE-S marker system will enable extension of plastid transformation to cereal crops.

The utility of the new chimeric promoters

The σ^{70} -type plastid ribosomal RNA operon promoter, Prrn, is the strongest known plastid promoter expressed in all tissue types. The ultimate product of this promoter in the plastid is RNA not protein. Therefore, a series of chimeric promoters were constructed to facilitate protein accumulation from Prrn, using expression of the neomycin phosphotransferase (NPTII) enzyme as the reference protein.

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- specific expression profiles. Some of the expression cassettes described here will facilitate relatively high levels of protein expression in all tissues, including leaves, roots and seeds. Other cassettes have different expression profiles: for example will facilitate moderate levels of protein accumulation in the leaves while lead to relatively high levels of protein accumulation in the roots. Accumulation of a protein at levels of 10% to 50% of total soluble protein is considered high-level protein expression; low-levels of protein expression would be in the range of ≤0.1% total soluble cellular protein.

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depends on the rate at which the gene product accumulates during the early stage of transformation. Since initially present only in a few copies per cell, high levels of expression from a few copies will provide protection from toxic substances early on, facilitating efficient recovery of transformed lines. The expression cassettes will be useful to drive the expression of the genes conferring resistance to the antibiotics

Efficiency of the selectable marker gene

streptomycin, spectinomycin and hygromycin, and the herbicides phosphinotrycin and glyphosate. In such applications addition of amino acids at the N-terminus is acceptable, as long as it does not interfere with the expression of the selectable marker genes. NPTII is such an enzyme. In cases like NPTII, an N-terminal fusion and thereby the mRNA "Downstream Box" sequences give an additional at least two to four-fold increase in protein levels. The -DB construct which relied on an NheI site, and involved addition of one (N-terminal) amino acid of the source gene coding region is convenient, but is not necessary. When translational fusion is not feasible due to inactivation of proteins, seamless in-frame constructs may be created by PCR methods outlined in the application.

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3) A second major area on which application of the chimeric promoters is extremely useful is protein expression for pharmaceutical, industrial or agronomic purposes. The examples include, but are not restricted to, production of vaccines, healthcare products like human hemoglobin, industrial or household enzymes.

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While certain of the preferred embodiments of the present invention have been described and specifically exemplified above, it is not intended that the invention be limited to such embodiments. Various modifications may be made thereto without departing from the scope and spirit of the present invention, as set forth in the following claims.

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What is claimed is:

1. A recombinant DNA construct for expressing at least one heterologous protein in the plastids of higher plants, said construct comprising a 5' regulatory region which includes a promoter element, a leader sequence and a downstream box element operably linked to a coding region of said at least one heterologous protein, said chimeric regulatory region enhancing translational efficiency of an mRNA molecule encoded by said DNA construct.

- 2. A vector comprising the DNA construct of claim 1.
- 3. A recombinant DNA construct as claimed in claim 1, said 5' regulatory region being selected from the group consisting of PrnnLatpB+DBwt, SEQ ID NO:1, PrrnLatpB-DB, SEQ ID NO:2, PrrnLatpB+DBm, SEQ ID NO:3, PrrnLclpP+DBwt, SEQ ID NO:4, PrrnclpP-DB, SEQ ID NO:5, PrrnLrbcL+DBwt, SEQ ID NO:6, PrrnLrbcL-DB, SEQ ID NO:7, PrrnLrbcL+DBm, SEQ ID NO:8, PrrnLpsbB+DBwt, SEQ ID NO:9, PrrnLpsbB-DB, SEQ ID NO:10, PrrnLpsbA+DBwt, SEQ ID NO:11, PrrnLpsbA-DB, SEQ ID NO:12, PrrnLpsbA-DB(+GC), SEQ ID NO:13.
- 30 4. A recombinant DNA construct as claimed in claim 1, said 5' regulatory region being selected from the group consisting of PrrnLT7g10+DB/Ec, SEQ ID NO:14,

PrrnLT7g10+DB/pt, SEQ ID NO:15, PrrnLT7g10-DB, SEQ ID NO:15.

5. A vector comprising a DNA construct as claimed in claim 1.

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- 6. A DNA construct as claimed in claim 1, said downstream box element having a sequence selected from the group consisting of 5'TCCAGTCACTAGCCCTTCGGCA'3 and 5'CCCAGTCATGAATCACAAAGTGGTAA'3.
- 7. A DNA construct as claimed in claim 1, wherein said heterologous protein is expressed from a bar gene encoded by S. hydroscopicus said bar gene inserted into a plasmid selected from the group consisting of pKO12, and pJEK3, said pJEK3 having the sequence of SEQ ID NO: 18.
- 8. A DNA construct as claimed in claim 1, wherein said heterologous protein is expressed from a synthetic bar encoding nucleic acid, said synthetic bar nucleic acid having selected from the group consisting of SEQ ID NO: 19 and SEQ ID NO:20.
 - 9. A DNA construct as claimed in claim 1, said at least one heterologous protein comprising a fusion protein.
- 30 10. A DNA construct as claimed in claim 9, said fusion protein having a first and second coding region operably linked to said 5' regulatory region such that production of said fusion protein is regulated by

said 5' regulatory region, said first coding region encoding a selectable marker gene and said second coding region encoding a fluorescent molecule to facilitate visualization of transformed plant cells.

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- 11. A vector comprising the DNA construct of claim 10.
- 12. A DNA construct as claimed in claim 9,
 said fusion protein consisting of an aadA coding region
 operably linked to a green fluorescent protein coding
 region.
- 13. A DNA construct as claimed in claim 10, said aadA coding region being operably linked to said green fluorescent protein coding region via a nucleic acid molecule encoding a peptide linker having a sequence selected from the group consisting of ELVEGKLELVEGLKVA and ELAVEGKLEVA.

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- 14. A DNA construct as claimed in claim 10, said construct having a sequence selected from the group of SEQ ID NOS: 21-25 and 27.
- 15. A plasmid for transforming the plastids of higher plants, said plasmid being selected from the group consisting of pHK30(B), pHK31(B), pHK60, pHK32(B), pHK33(B), pHK34(A), pHK35(A), pHK64(A), pHK36(A), pHK37(A), pHK38(A), pHK39(A), pHK40(A), pHK41(A), pHK42(A), pHK43(A), pMSK56, pMSK57, pMSK48, pMSK49, pMSK35, pMSK53 and pMSK54.

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16. A transgenic plant containing a plasmid as claimed in claim 15.

- 17. A transgenic plant as claimed in claim15, said plant being selected from the group consisting of monocots and dicots.
 - 18. A method for producing transplastomic monocots, comprising:
 - a) obtaining embryogenic cells;

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- b) exposing said cells to a heterologous DNA molecule under conditions whereby said DNA enters the plastids of said cells, said heterologous DNA molecule encoding at least one exogenous protein, said at least one exogenous protein encoding a selectable marker;
- c) applying a selection agent to said cells to facilitate sorting of untransformed plastids from transformed plastids, said cells containing transformed plastids surviving and dividing in the presence of said selection agent;
- d) transferring said surviving cells to selective media to promote shoot regeneration and growth; and
- e) rooting said shoots, thereby producingtransplastomic monocot plants.
 - 19. A method as claimed in claim 18, wherein said heterologous DNA molecule is introduced into said plant cell via a process selected from the group consisting of biolistic bombardment, Agrobacterium- mediated transformation, microinjection and electroporation.

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20. A method as claimed in claim 18, wherein protoplasts are obtained from said embryogenic cells and said heterologous DNA molecule is delivered to said protoplasts by exposure to polyethylene glycol.

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21. A method as claimed in claim 18, wherein said selection agent is selected from the group consisting of streptomycin, and paromomycin

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- 22. A monocot transformed via the method of claim 18.
- 23. A transformed monocot plant as claimed in claim 22, said monocot plant being selected from the group consisting of maize, millet, sorghum, sugar cane, rice, wheat, barley, oat, rye, and turf grass.
- 24. A method for producing transplastomic rice20 plants, said method comprising:
 - a) obtaining embryogenic calli;
 - b) inducing proliferation of calli on modified CIM medium;
 - c) obtaining embryogenic cell suspensions of said proliferating calli in liquid AA medium;
 - d) bombarding said embryogenic cells with microprojectiles coated with plasmid DNA;
 - e) tranferring said bombarded cells to selective liquid AA medium;
 - f) transferring said cells surviving in AA medium to selective RRM regeneration medium for a time period sufficient for green shoots to appear; and

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 $\ensuremath{\mathtt{g}})$ rooting said shoots in a selective MS salt medium.

- DNA being selected from the group of plasmids consisting of pMSK35 and pMSK53, pMSK54 and pMSK49.
 - 26. A transplastomic rice plant produced by the method of claim 24.

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- 27. A method for containing transgenes in transformed plants, comprising:
- a) determining the codon usage in said plant
 to be transformed and in microbes found in association with said plant; and
 - b) genetically engineering said transgene sequence via the introduction of rare codons to abrogate expression of said transgene in said plant associated microbe.
 - 28. A method as claimed in claim 27, wherein said transgene is a bar gene and said rare codons are arginine encoding codons selected from the group consisting of AGA and AGG.

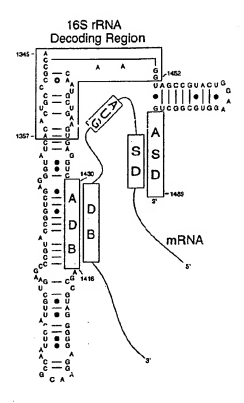


Figure 1A

1 10 20 26

pt ADB 3'-AGGUCAGUGAUCGGGACGGAAGCCGU-5'
1430 1416

1 10 20 26

Ec ADB 3'-GGGUCAGUACUUAGUGUUUCACCAUU-5'
1483 1469

Figure 1B

```
atpB wild type
   mRNA -
             AUGAGAAUCAAUCCUACUACUUCUGGUUCUGGGGUUUCCACGCUUGAAAA
              3'-AGGUCAGUSAUCGGGACGGAAGCCGU-5' (9/7)
           - NGGAGANUANACCCGACAACAAGUGGAAGUGGGGUGUCCACGGCUAGC
                       -AGGUCAGUGAUCGGGACGGAAGCCGU-5 (11/9)
              3 - AGGUCAGUGAUCGGGACGGAGCCGU-5' (10/8)
         3'-AGGUCAGUGAUCGGGACGGAAGCCGU-5' (14)

***OGGUCAGUGAUCGGGACGGAAGCCGU-5' (14)

***OGGUCAGUGAUCGGGACGGAAGCCGU-5' (14)

***DGCCUAUUGGUGAUCGGACGGAAGCCGU-5' (14)

***DGCCUAUUGGUGAUCGGACGGAAGCCGU-5' (14)

***OGGUCAGUGAUCGGGACGGAAGCCGU-5' (13)
clpP wild type
   mRNA -
            3'-AGGUC<u>AGUGAUCGGGACGGA</u>AGCCGU-5' (13)
           rbcL wild type
   mRNA .
                       3'-AGGUCAGUGAUCGGGACGGAAGCCGU-5' (10/5)
           3'-AGGUCAGUGAUCGGACGGAAGCCGU-5' (9/5)
          psbB wild type
   mRNA -
psbA wild type
          mRNA -
```

Figure 2A

Figure 2B

PrrnLatpB+DBwt (pHK10)

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG AATTAACCGA
- 101 TCGACGTGCa AGCGGACATT TATTTTAAAT TCGATAATTT TTGCAAAAAC
- 151 ATTTCGACAT ATTTATTTAT TTTATTATTA TGAGAATCAA TCCTACTACT NheT
- 201 TCTGGTTCTG GGGTTTCCAC Ggctagc

PrrnLatpB-DB (pHK11)

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGCCTATA TTTCTGGGAG AATTAACCGA
- 101 TCGACGTGCa AGCGGACATT TATTTTaAAT TCGATAATTT TTGCAAAAAC
- .151 ATTTCGACAT ATTTATTTAT TTTATTATTA TGAGAgctag c

PrinLatpB+DBm (pHK50)

- SacI
 1 gagete<u>GCTC CCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG</u>
- 51 GATTGACGIG AGGGGCAGG GATGGCTATA TTTCTGGGAG AATTAACCGA
- 101 TCGACGTGCA AGCGGACATT TATTTTAAAT TCGATAATTT TTGCAAAAAC
- 151 ATTTCGACAT ATTTATTTAT TTTATTATTA TGAGAATAAA cCCgACaACa
- 201 agTGGaagTG GGGTgTCCAC Ggctagc

PrrnLclpP+DBwt (pHK12)

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG TTACGTTTCC
 - 101 ACCTCAAAGT GAAATATAGT ATTTAGTTCT TTCTTTCATT TAATGCCTAT
 - 151 TGGTGTTCCA AAAGTCCCTT TCCGAAGTCC TGGAGAGGAA gctage

PrrnLclpP-DB (pHK13)

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG TTACGTTTCC
- 101 ACCTCAAAGT GAAATATAGT ATTTAGTTCT TTCTTTCATT TAATGCCTgc
- 151 tage

Figure 3A

PrrnLrbcL+DBwt (pHK14)

SacI

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG TCGAGTAGAC
- 101 CTTGTTGTTG TGAaAATTCT TAATTCATGA GTTGTAGGGA GGGATTTATG
 NheI
- 151 TCACCACAAA CAGAGACTAA AGCAAGTGTT GGATTCAAAg ctagc

PrrnLrbcL-DB (pHK15)

SacI

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG TCGAGTAGAC
- 101 CTTGTTGTTG TGAaAATTCT TAATTCATGA GTTGTAGGGA GGGATTT**ATG**NheI
- 151 TCAgctagc

PrrnLrbcL+DBm (pHK54)

SacI

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG TCGAGTAGAC
- 101 CTTGTTG TGABARTTCT TARTTCATGA GTTGTAGGGA GGGATTTATG
- 151 aguCCuCAgA CAGAaACaAA AGCcucaGTa GGATTCAAAg ctagc

PrrnLpsbB+DBwt (pHK16)

SacI

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG CAATGCAATA
- 101 AAGTTACGTA GTGTCTATTT ATCTTTGATA TAAGGGGTAT TTCCATGGGT
 - Nhel
- 151 TTGCCTTGGT ATCGTGTTCA TACCGTTGTA TTGAATGATg ctagc

PrrnLpsbB-DB (pHK17)

SacI

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG CAATGCAATA
 - Ncol Nhel
- 101 AAGTTACGTA GTGTCTATTT ATCTTTGATA TAAGGGGTAT TTccatggct
- 151 agc

Figure 3B

PrrnLpsbA+DBwt (pHK21)

- SacI
 1 gagete<u>GCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG</u>
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAA AAAAGCCTTC
- 101 CATTTCTAT TTTGATTTGT AGAAAACTAG TGTGCTTGGG AGTCCCTGAT
- 151 GATTAAATAA ACCAAGATTT TACCATGACT GCAATTTTAG AGAGAGCtag
- 201 c

PrrnLpsbA-DB (pHR22)

SacI

- 1 gagctcGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAA AAAAGCCTTC
- 101 CATTTTCTAT TTTGATTTGT AGAAAACTAG TGTGCTTGGG AGTCCCTGAT Ncol Nhel
- 151 GATTAAATAA ACCAAGATTT TAccatggct agc

PrrnLpsbA-DB(+GC) (pHK23)

SacI

- 1 gagctcGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG CAAAAAGCCT
- 101 TCCATTTTCT ATTTTGATTT GTAGAAAACT AGTGTGCTTG GGAGTCCCTG Ncol Nhel
- 151 ATGATTAAAT AAACCAAGAT TTTAccatgg ctagc

Figure 3C

PrrnLT7g10+DB/Ec (pHK18)

SacI

- 1 gageteGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG GGAGACCACA
- 101 ACGGTTTCCC aCTAGAAATA ATTTTGTTTA ACTTTAAGAA GGAGATATAC NheI
- 151 ATATGGCaAG CATGACTGGT GGACAGgcta gc

PrrnLT7g10+DB/pt (pHK19)

SacI

- 1 gagctcGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG GGAGACCACA
- 101 ACGGTTTCCC aCTAGAAATA ATTTTGTTTA ACTTTAAGAA GGAGATATAC NheI
- 151 ATATGGCaAt cactageect geettGgeta ge

PrrnLT7g10-DB (pHK20)

- SecI
- 1 gagctcCCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG AGGCTCGTGG
- 51 GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG GGAGACCACA
 - 101 ACGGTTTCCC aCTAGAAATA ATTTTGTTTA ACTTTAAGAA GGAGATATAC . NheI
 - 151 ATATGgctag c

Figure 3D

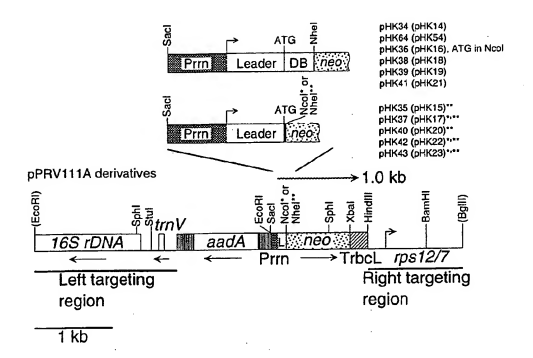


Figure 4A

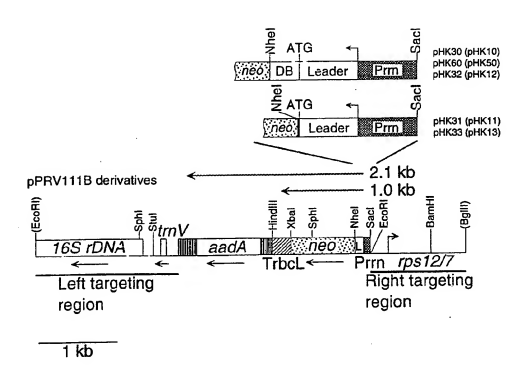
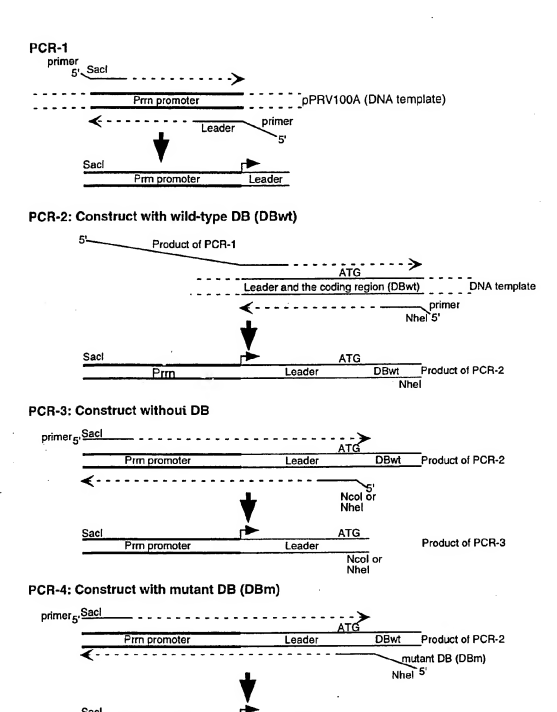


Figure 4B

Product of PCR-4



Leader

Figure 5

Prm promoter

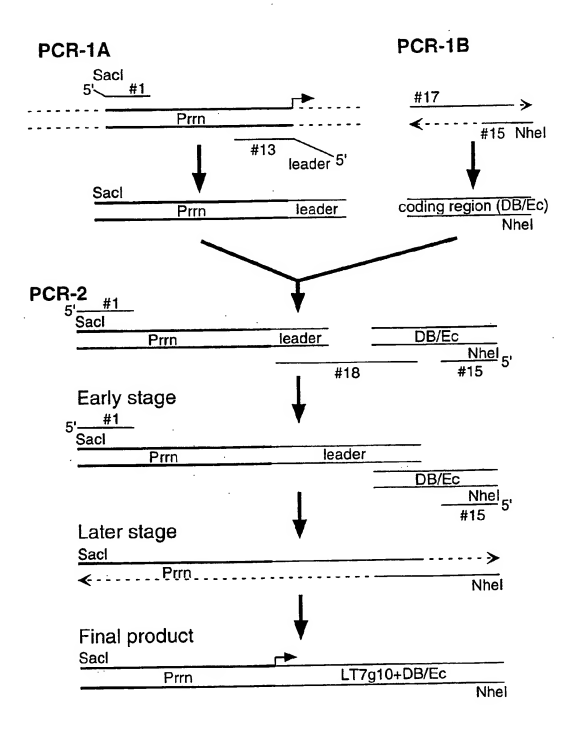


Figure 6

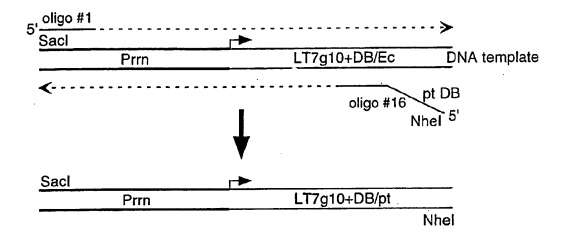


Figure 7

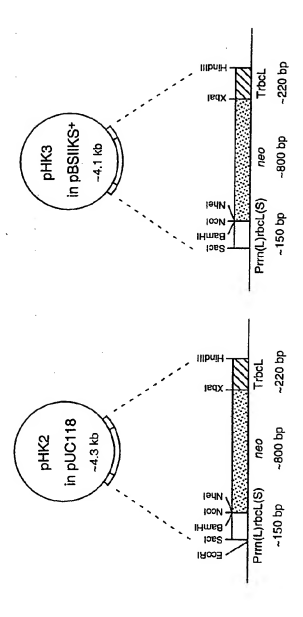
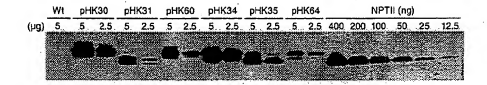
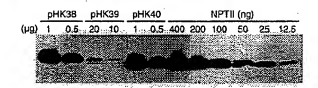
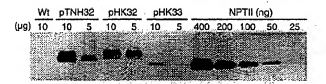


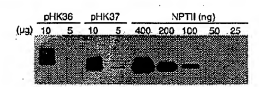
Figure 8

	SacI	
1	gageteggta eccaaaGCTC CCCCGCCGTC GTTCAATGAG AATGGATAAG	
51	AGGCTCGTGG GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG	Ι
101	CGAACTCCGG GCGAATACGA AGCGCtTGGA TACAGTTGTA GGGAGGGATC NheI	
151	catggctagc ATTGAACAAG ATGGATTGCA CGCAGGTTCT CCGGCCGCTT	
201	GGGTGGAGAG GCTATTCGGC TATGACTGGG CACAACAGAC AATCGGCTGC	
251	TCTGATGCCG CCGTGTTCCG GCTGTCAGCG CAGGGGCGCC CGGTTCTTTT	
301	TGTCAAGACC GACCTGTCCG GTGCCCTGAA TGAACTCCAG GACGAGGCAG	
351	CGCGGCTATC GTGGCTGGCC ACGACGGGCG TTCCTTGCGC AGCTGTGCTC	
401	GACGTTGTCA CTGAAGCGGG AAGGGACTGG CTGCTATTGG GCGAAGTGCC	
451	GGGGCAGGAT CTCCTGTCAT CTCACCTTGC TCCTGCCGAG AAAGTATCCA	
501	TCATGGCTGA TGCAATGCGG CGGCTGCATA CGCTTGATCC GGCTACCTGC	
551	CCATTCGACC ACCAAGCGAA ACATCGCATC GAGCGAGCAC GTACTCGGAT	
601	GGAAGCCGGT CTTGTCGATC AGGATGATCT GGACGAAGAG CATCAGGGGC	
651	TCGCGCCAGC CGAACTGTTC GCCAGGCTCA AGGCGCGCAT GCCCGACGGC	
701	GAGGATCTCG TCGTGACACA TGGCGATGCC TGCTTGCCGA ATATCATGGT	
751	GGAAAATGGC CGCTTTTCTG GATTCATCGA CTGTGGCCGG CTGGGTGTGG	
801	CGGACCGCTA TCAGGACATA GCGTTGGCTA CCCGTGATAT TGCTGAAGAG	
851	CTTGGCGGCG AATGGGCTGA CCGCTTCCTC GTGCTTTACG GTATCGCCGC	
901	TCCCGATTCG CAGCGCATCG CCTTCTATCG CCTTCTTGAC GAGTTCTTCT XbaI	
951	GAgcgggtct agagtAGACA TTAGCAGATA AATTAGCAGG AAATAAAGAA	
1001	GGATAAGGAG AAAGAACTCA AGTAATTATC CTTCGTTCTC TTAATTGAAT	
1051	TGCAATTAAA CTCGGCCCAA TCTTTTACTA AAAGGATTGA GCCGAATACA	
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1151	TACAAGATTT GAAATACAAA ATCTAGcaag ctt	









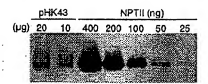


Figure 10

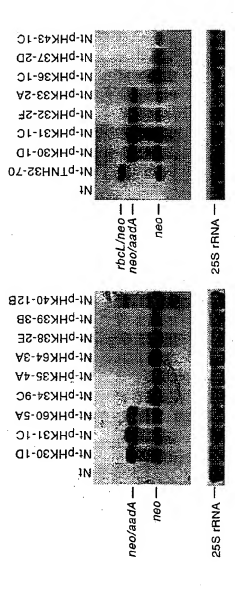
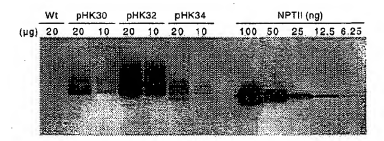


Figure 11

### WE AND AGA AND CAN AND CAN UCU GGD UUT GGG GUU UCC ACG Fraction 1.0 0.22 0.27 0.61 0.30 0.37 0.37 0.38 0.31 0.38 0.31 0.26 0.35 0.14 0.15 atpliet/1000 24.6 7.8 15.5 18.1 13.5 18.4 18.4 20.2 28.2 20.2 19.2 24.9 9.1 7.5 atpliet/1000 24.6 7.8 15.5 18.1 13.5 18.4 18.4 20.2 28.2 20.2 19.2 24.9 9.1 7.5 atpliet/1000 24.6 7.8 15.5 18.1 13.5 18.4 18.4 20.2 28.2 20.2 19.2 24.9 9.1 7.5 atpliet/1000 24.6 1.2 10.2 0.29 0.30 0.30 0.30 0.30 0.30 0.30 0.30 0.3	111111111111111111111111111111111111111															
1.0 0.22 0.27 0.61 0.30 0.37 0.37 0.31 0.38 0.31 0.26 0.35 0.14 AUG AGA AUA AAC CCG ACA ACA a gU Gga agU GGG GUG UCC Net Arg Ille Asn Pro Thr Thr Ser Gly Ser Gly Val Ser Ill 0.22 0.29 0.39 0.30 0.23 0.23 0.14 0.24 0.14 0.26 0.21 0.21 0.14 AUG UCA CCA CAA ACA GAG ACU AAA GCA AGU GUU GGA UUC Ser Gly Thr GlU Thr GlU Thr Ill 9.3 19.2 15.3 9.1 AUG UCA CCA CAA ACA GAG ACU AAA GCA AGU GUU GGA UUC Ser Gly Thr GlU Thr GlU Thr Ill 9.3 19.2 15.3 9.1 AUG UCA CCA CAA ACA GAG ACU AAA GCA AGU GUU GGA UUC Ser Gly Thr GlU Thr Ill 9.3 19.2 15.3 9.1 AUG UCA CCA CAA ACA GAG ACU AAA GCA AGU GUU GGA UUC Ser Gly Thr GlU Thr Ill 9.3 19.2 15.3 9.1 AUG Ser Pro Gln Thr Glu Thr Ill 9.3 24.9 17.9 22.5 AUG GU CCU CAG ACA GAG ACA ACA AAA GCC uca GUU GGA UUC Ser Pro Gln Thr Glu Thr Ill 9.3 24.9 17.9 22.5 AUG GU GAJ D.24 0.57 0.23 0.52 0.23 0.60 0.18 0.11 0.24 0.40 0.40 24.6 13.5 10.6 21.0 11.7 12.4 18.4 22.0 18.1 9.3 24.9 17.9 22.5 AUG GCA ACA ACA ACA ACA ACA AAA GCC uca GUU GGA UCC Met Ser Pro Gln Thr Glu Thr Ill 22.0 10.1 13.5 21.8 17.9 22.5 AUG GCA AGC AUG ACU GGU GGA CAG GCU agc auu gaa caa Met Ala Ser Met Thr Gly Gly Gln Ala Ser Ille Glu Gln 1.0 0.29 0.07 1.00 0.37 0.38 0.24 0.43 0.39 0.07 0.45 0.62 0.57 0.40 AUG GCA AUC AUG ACU GGU GGA CAG GCU agc auu gaa caa Met Ala Ser Met Thr Gly Gly Gln Ala Ser Ille Glu Gln 1.0 0.29 0.07 1.00 0.37 0.38 0.24 0.43 0.39 0.07 0.45 0.62 0.57 0.40 0.29 0.07 0.45 0.62 0.57 0.75 0.24 0.40 0.29 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.07 0.45 0.62 0.57 0.75 0.20 0.20 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 0.39 0.07 0.45 0.62 0.57 0.75 0.20 0.20 0.20 0.20 0.20 0.27 0.37 0.37 0.37 0.34 0.34 0.34 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 0.30 0.30 0.30 0.30 0.30 0.30 0.30	acps we	AUG Met				CCU	ACU	ACU		GGU	מכמ	966			ACG	
AUG AUG AUG ACG C12 C22 C24 C11 T11 T11 T24 18.4 C2.0 C13 C13 C14 C15 C15 C17 C17 </td <td>Fraction Triplet/1000</td> <td>1.0</td> <td></td> <td></td> <td></td> <td>0.30</td> <td>0.37.</td> <td>0.37</td> <td></td> <td>0.38 28.2</td> <td>0.31 20.2</td> <td>61Y 0.26 19.2</td> <td></td> <td></td> <td>Thr 0.15 7.5</td> <td></td>	Fraction Triplet/1000	1.0				0.30	0.37.	0.37		0.38 28.2	0.31 20.2	61Y 0.26 19.2			Thr 0.15 7.5	
AUG CCA CAA ACA CAC ACA CAC ACA ACA ACA CAC AAA GCA AGU GUU GGA UUC 1.0 0.21 0.24 0.57 0.23 0.38 0.37 0.60 0.29 0.14 0.35 0.24 0.40 2.4.6 13.5 10.6 21.0 11.7 12.4 18.4 22.0 18.1 9.3 24.9 17.9 22.5 AUG GU CAG GAA ACA AAA GCC uca GU GGA UUC Met Ser Pro GIN Thr GLU Thr LVS AAA GCA UUC GGA UUC UUC AUG ACA GAA ACA AAA GCA UUC AUG DUC UUC AUG ACA AAA ACA UUC AUG CU CU DUC DUC DUC DUC DUC DUC	atpB m Fraction Triplet/1000	AUG Met 1.0 24.6			AAc Asn 0.39 11.4			ACa Thr 0.23 11.7	agu Ser 0.14 9.3	Gga Gly 0.24 17.9	agu Ser 0.14 9.3	GGG Gly 0.26 19.2	GUG Val 0.21 15.3	UCC Ser 0.14 9.1	ACG Thr 0.15 7.5	
AUG GCA AUG ACU GGU GGA CAG GCA GCA GCA GCA GCA GCA GCA GCA	rbcL wt	AUG Met	UCA	CCA	CAA		GAG			GCA	AGU			COC	AAA	
AUG agu CCu CAg ACA GAA ACA AAA GCc uca GUa GGA UUC 1.0 0.14 0.30 0.43 0.23 0.62 0.23 0.60 0.16 0.21 0.31 0.24 0.40 24.6 9.3 13.5 15.5 11.7 20.7 11.7 22.0 10.1 13.5 21.8 17.9 22.5 AUG GCa AUG GGU GGA CAG gcu agc auu gaa caa 1.0 0.29 0.07 1.00 0.37 0.38 0.24 0.43 0.39 0.07 0.42 0.52 24.6 18.1 4.7 24.6 18.4 28.2 17.9 15.5 24.4 4.7 25.9 20.7 21.0 AUG GCa AUG GGA CCu GCA 0.43 0.39 0.07 0.45 0.65 0.57 21.0	Fraction Triplet/1000	1.0	0.21	0.24	0.57 21.0		0.38 12.4			Ala 0.29 18.1	Ser 0.14 9.3			Phe 0.40 22.5	Lys 0.60 22.0	
1.0 0.14 0.30 0.43 0.23 0.62 0.23 0.60 0.16 0.21 0.31 0.24 0.40 24.6 9.3 13.5 15.5 11.7 20.7 11.7 22.0 10.1 13.5 21.8 17.9 22.5 AUG GCa AGC AUG ACU GGU GGA CAG gcu agc auu gaa caa Met Ala Ser Met Thr Gly Gly Gln Ala Ser Ile Glu Gln 1.0 0.29 0.07 1.00 0.37 0.38 0.24 0.43 0.39 0.07 0.45 0.62 0.57 24.6 18.1 4.7 24.6 18.4 28.2 17.9 15.5 24.4 4.7 25.9 20.7 21.0 AUG GCa Auc agc ccu gcc uuG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 21.0 AUG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 21.0 AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Mct Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	rbcL m	AUG Met	agu Ser	CCu	CAg									UUC	AAA	
AUG GCa AGC AUG ACU GGU GGA CAG gcu agc auu gaa caa Met Ala Ser Met Thr Gly Gly Gln Ala Ser Ile Glu Gln 1.0 0.29 0.07 1.00 0.37 0.38 0.24 0.43 0.39 0.07 0.45 0.62 0.57 24.6 18.1 4.7 24.6 18.4 28.2 17.9 15.5 24.4 4.7 25.9 20.7 21.0 AUG GCa Auc acu agc ccu gcc uuG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 24.6 18.1 15.5 18.4 4.7 13.5 10.1 34.7 24.4 4.7 25.9 20.7 21.0 AUG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 24.6 18.1 15.5 18.4 4.7 13.5 10.1 34.7 24.4 4.7 25.9 20.7 21.0 AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Mct Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	Fraction	1.0	0.14	0.30	0.43									Phe	Lys	
AUG GCa AGC AUG ACU GGU GGA CAG gcu agc auu gaa caa 1.0 0.29 0.07 1.00 0.39 0.24 0.43 0.39 0.07 0.45 0.62 0.57 24.6 18.1 4.7 24.6 18.4 28.2 17.9 15.5 24.4 4.7 25.9 20.7 21.0 AUG GCa Auc acu agc ccu gcc uuG gcu agc auu gaa caa 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 21.0 24.6 18.1 15.5 18.4 4.7 13.5 10.1 34.7 24.4 4.7 25.9 20.7 21.0 24.6 18.1 15.5 18.4 4.7 13.5 10.1 34.7 24.4 4.7 25.9 <td>Triplet/1000</td> <td>24.6</td> <td>9.3</td> <td>13.5</td> <td>15.5</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>22.5</td> <td>22.0</td> <td></td>	Triplet/1000	24.6	9.3	13.5	15.5									22.5	22.0	
1.0 0.29 0.07 1.00 0.37 0.38 0.24 0.43 0.39 0.07 0.45 0.62 0.57 24.6 18.1 4.7 24.6 18.4 28.2 17.9 15.5 24.4 4.7 25.9 20.7 21.0 AUG GCa Auc acu agc ccu gcc uuG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 4.7 25.9 20.7 21.0 AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Mct Ala Ser Ile Glu Gln 34.7 24.4 4.7 25.9 20.7 21.0 AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Mct Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 20.2 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	T7g10+DB/Ec	AUG	GCa	AGC										caa	gan	
AUG GCa Auc acu agc ccu gcc uuG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 21.0 Aug gcu agc auu gaa caa gau gaa uug cac gca ggu ucu Mct Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	Fraction	1.0	0.29	0.07										GIn	Asp 0 75	
AUG GCa Auc acu agc ccu gcc uuG gcu agc auu gaa caa Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 24.6 18.1 15.5 18.4 4.7 13.5 10.1 34.7 24.4 4.7 25.9 20.7 21.0 AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Met Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	Triplet/1000	24.6	18.1	4.7										21.0	24.6	
Met Ala Ile Thr Ser Pro Ala Leu Ala Ser Ile Glu Gln 1.0 0.29 0.27 0.37 0.07 0.30 0.16 0.24 0.39 0.07 0.45 0.62 0.57 24.6 18.1 15.5 18.4 4.7 13.5 10.1 34.7 24.4 4.7 25.9 20.7 21.0 AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Mct Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	T7g10+DB/pt	AUG			acn	agc .	ככת				aga	ลนน	gaa	Gas	ແອກ	
AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Met Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.55 20.7 21.0 24.6 24.4 4.7 25.9 20.7 21.0 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	Fraction	Met			Thr	Ser	Pro				Ser	Ile	Glu	Gln	Asp	
AUG gcu agc auu gaa caa gau gga uug cac gca ggu ucu Met Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	Triplet/1000	24.6			18.4	4.7	13.5				4.7	0.45	0.62	0.57	0.75 24.6	
Mct Ala Ser Ile Glu Gln Asp Gly Leu His Ala Gly Ser 1.0 0.39 0.07 0.45 0.62 0.57 0.75 0.24 0.24 0.28 0.29 0.38 0.31 24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	T7g10-DB	AUG					caa	gan	gga	bnr	gac	800			טט	
24.6 24.4 4.7 25.9 20.7 21.0 24.6 17.9 34.7 9.1 18.1 28.2 20.2	Fraction	Met					Gln	Asp (31y 1	ren 1	His	Ala	31y	Ser	Pro	
	Triplet/1000	24.6					21.0	24.6	17.9).24 (34.7	9.1	0.29 (18.1 ;	0.38 (0.31	0.30	

Figure 12



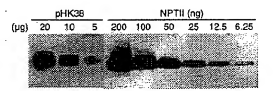


Figure 13A

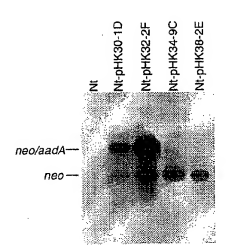


Figure 13B

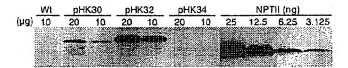


Figure 14

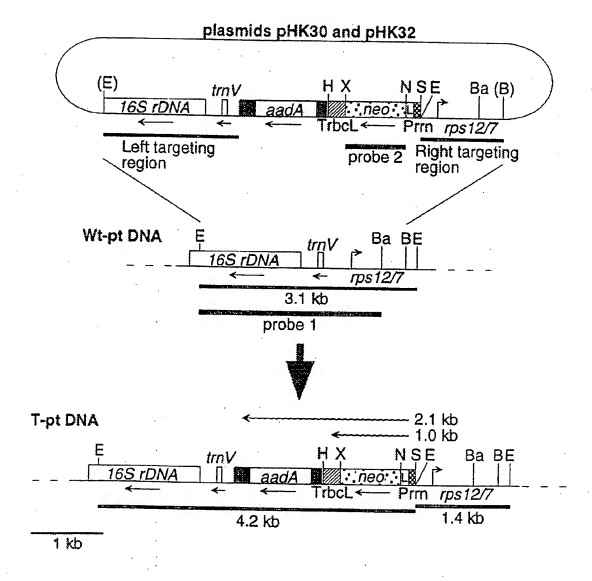


Figure 15A

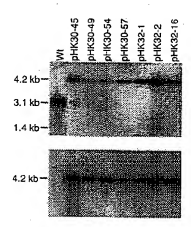


Figure 15B

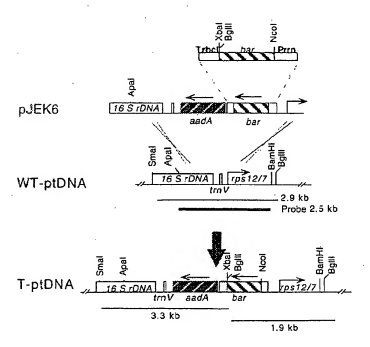


Figure 16A

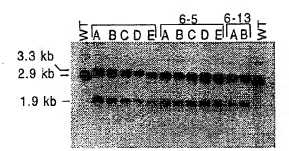


Figure 16B

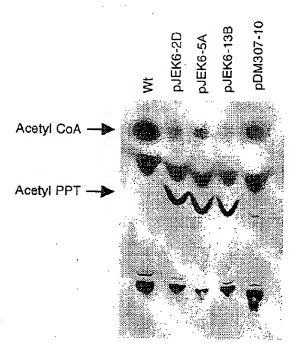
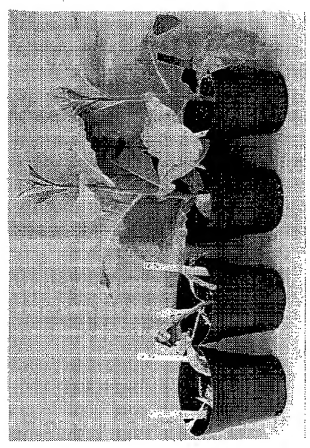


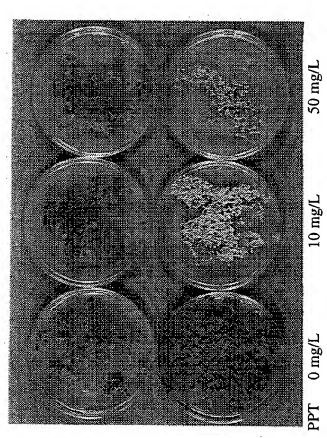
Figure 17



-6-2D Nt-6-5A Nt-wt Nt-wt

Figure 18A





W Q x pJEK6-5A O pJEK6-5A Q x wt o

NCO.	rGq	cac	caca	aaa	caga	agĄ(GCC	CAG	AAC	GAC	GCC	CGG	CCG	ACA'	TCC	GCC	GTĠ	CCA	ÇCG	
GGT2	ACc	gtg	gtgi	ttt	gtct E	CTC	CGGG	GTC'	TTG	CTG	CGG	GCC	GGC'	FGT	AGG	CGG	CAC	GGT (GC	60
AGG	CGG	ACA	rgc(CGG	CGGT	CTC	GCAC	CCA:	rcg'	FCA	ACC!	ACTA	ACA:	rcg/	AGA	CAA	GCA(GGI	CA	
TCC	GCC:	rgt <i>i</i>	ACGO	GCCC	GCCA V	GAC	CGT	GT!	AGC	AGT	rgg7	[GA]	rgt <i>i</i>	AGC T	rcT(-+- GTT(CGT(GCCA	+ \GT	120
										N		Y	-	_	T	S	_	V		
ACT1						+			+-	·		+				-+			-+	180
TGA	AGG(R	T T	GCT E	CGG P	CGT Q	'CCT E	TGC P	Q Q	rcc1 E	CAC W	CTC T	CCT D	rgci D	'GG <i>F</i> L	AGC <i>I</i> V	AGG(R	CAGA L	ACGC R		
AGC	CTA	TCC	CTG	GCI	CGT	CGC	CGA	GGT	GGA	ACGO	CGA	GGT	'CGC	CGG	CAT	CGC	CTA	CGC	GG	0.40
TCGC	GAT	'AGG P	GAC	CGA	GCA V	GCG	GCT	'CCA	CCI	GCC	GCT	CCA	IGC G	GCC	GTA	\GC@	GAT Y	'GCG	CC G	240
GCCC	СТС	GAA	.GGC	ACG	CAA	CGC	СТА	CGA	CTC	GAC	GGC	CGA	GTC	GAC	:CG'I	'GTA	CGT	CTC	CC	
CGGG	GAC	CTT	CCG	TGC	 GTT N	GCG	GAT	GCT	'GAC	CTG	CCG	GCT	CAG	CTG	GCA	САТ	CCA	GAG	GG	300
CCCG	CCA	.CCA	.GCG	GAC	GGG.	ACT	GGG	CTC	CAC	GCT	CTA	CAC	CCA	CCT	GCT	GAA	.GTC	ССТ	GG	
GGGC	GGT	GGT	CGC	CTG	CCC.	TGA	ccc	GAG	GTG	CGA	GAT	GTG	GGT	GGA	.CGA	CTT	CAG	GGA	-+ CC	360
		Q			G														E	
AGGC		+				+			-+-			+				+			-+	420
TCCG	TGT	CCC	GAA	GTT	CTC(S	GCA(CCA	GCG.	ACA	GT.A	GCC	CGA	CGG	GTT	GCT	GGG P	CTC	GCA(CG R	720
GCAT	GCA	CGA	GGC	GCT	CGG	ATA	rgc	CCC	CCG	CGG	CAT	GCT	GCG	GGC	GGC	CGG	CTT	CAA	GC	
CGTA	CGT	GCT(CCG	CGA	G G	rat <i>i</i>	ACG	GGG	GGC	GCC	GTA	CGA	CGC	CCG	CCG	GCC	GAA	GTT	-+ CG H	480
ACGG(GAA	CTG	GCA:	rga(CGTC	GGT	TTT	CTG	GCA	GCT	GGA	CTT	CAG	CCT	GCC	GGT	ACC	GCC	cc	
rGCC	CTT	3AC(CGTA	ACT(GCAC	CCC)AA/	GAC	CGT	CGA	CCT	GAAG	STC	GGA	CGGC	CA	rgga	cec	:0	540
G	14	VV	п		ν.		r glI		Q	L	ט	F'	S	L	P	V	P	P	R	
GTCC	GGT	CCT	GCC	CGTC	CAÇC	GAG	ATC	CTGA	ATG?	Atc	gaat	tco	etge	cago	ccc	gggg	ggat	icca	ıc	
CAGG	CCA	GGAC	CGGC	CAC		CTC	TAG		-+			+-			1				+ 4	600
agt	Xba	iga	610												,					
tcaa			OIC	,						1	Fi	gu	re	1	19					

Ncol Nhel

<u>CcATGq</u>ctAGCCCAGAAaGAaGaCCGGCCGAtATtaGaCGTGCtACaGAaGCtGAtATGC ggTACcgaTCGGGTCTTtCTtCtGGCCGGCTaTAatCtGCACGaTGtCTtCGaCTaTACG M A S P E R R P A D I R R A T E A D M P CaGCaGTtTGtACaATtGTtAAtCAtTAtATaGAaACAAGtACcGTaAACTTtcGaACtG A V C T I V N H Y I E T S T V N F R T E ${\tt AaCCtCAaGAACCtCAaGAaTGGACtGAtgAttTaGTCCGTtTaCGaGAGCGCTATCCtT}$ ------ ${\tt TtGGaGTtCTTGGaGTtCTtACCTGaCTaCTaaAtCAGGCAaAtGCtCTCGCGATAGGaA}$ PQEPQEWTDDLVRLRERYPW ${\tt GGCTtGTaGCaGAaGTtGACGGaGAaGTaGCtGGgATtGCaTAtGCGGGCCCgTGGAAaG}$ ${\tt CCGAaCAtCGtCTtCAaCTGCCtCTtCAtCGaCCcTAaCGtATaCGCCCGGGcACCTTtC}$ L V A E V D'G E V A G I A Y A G P W K A ${\tt CAcGaAAtGCaTAtGAtTGGACgGCtGAaTCaACtGTgTACGTtTCaCCaCGtCAtCAaC}$ -------GTqCtTTaCGtATaCTaACCTGcCGaCTtAGtTGaCAcATGCAaAGtGGtGCaGTaGTtG R N A Y D W T A E S T V Y V S P R H Q R ${\tt GqACaGGACTtGGtTCtACttTaTAtACcCAtCTaCTGAAaTCttTGGAGGCACAgGGtT}$ -----+----+----+----+----+----+ ${\tt CcTGtCCTGAaCCaAGaTGaaAtATaTGgGTaGAtGACTTtAGaaACCTCCGTGTcCCaA}$ T G L G S T L Y T H L L K S L E A Q G F TtAAGAGtGTgGTaGCTGTtATaGGatTGCCgAAtGAtCCctcgGTaCGCATGCAcGAaG AaTTCTCaCAcCAtCGACAa'TAtCCtaACGGcTTaCTaGGgagcCAtGCGTACGTgCTtC K S V V A V I G L P N D P S V R M H E A CtCTcGGATATGCtCCcaGaGGtATGtTGaGGGCcGCaGGtTTCAAaCAtGGaAAtTGGC ------ ${\tt GaGAgCCTATACGaGGgtCtCCaTACaACtCCCGgCGtCCaAAGTTtGTaCCtTTaACCG}$ L G Y A P R G M L R A A G F K H G N W H TACTaCAtCCAAAaACCGTtGAaCTgAAGagaaAtGGtCATGGaGGaGCAGGgCAaaAtG D V G F W Q L D F S L P V P P R P V L P

BglII XbaI
CcGTtACtGAGATCTGATGAtctaga
-----+----GgCAaTGaCTCTAGACTACTagatct
V T E I * *

Figure 20A

NCOI NheI ccatgctagcccagaaagaacccgccgatattagacgtgctacagaagctgatatgc ______ ggTACcgaTCGGGTCTTtCTtCtGGCCGGCTaTAatCtGCACGaTGtCTtCGaCTaTACG MASPERRPADIRRATEADMP CaGCaGTtTGtACaATtGTtAAtCAtTAtATaGAaACAAGtACaGTaAAtTTtcGaACtG GtCGtCAaACaTGtTAaCAaTTaGTaATaTAtCTtTGTTCaTGtCAtTTaAAagCtTGaC A V C T I V N H Y I E T S T V N F R T E AaCCtCAaGAACCtCAaGAaTGGACtGAtGAttTaGTaCGTtTaCGaGAaCGtTATCCtT ______ TtGGaGTtCTTGGaGTtCTtACCTGaCTaCTaaAtCAtGCAeAtGCtCTtGCaATAGGaA PQEPQEWTDDLVRLRERYPW GGCTtGTaGCaGAaGTtGAcGGaGAaGTaGCtGGaATtGCaTAtGCtGGtcCgTgGAAaG ------ ${\tt CCGAaCAtCGtCTtCAaCTgCCtCTtCAtCGaCCtTAaCGtATaCGaCCaGGcACCTTtC}$ L V A E V D G E V A G I A Y A G P W K A CAcGaAAtGCaTAtGAtTGGACaGCtGAaTCaACtGTtTAtGTtTCaCCaCGtCAtCAaC GTGCtTFaCGtATaCTaACCTGtCCaCTtAGtTGaCAaATaCAaAGtGGtGCaGTaGTtG RNAYDWTAESTVYVSPRHQR GtaCaGGACTtGGtTCtACttTaTAtACtCAtCTtCTtAAaTCttTGGAaGCACAaGGtT CaTGtCCTGAaCCaAGaTGaaAtATaTGaGTaGAaGAaTTtAGaaACCTtCGTGTtCCaA TGLGSTLYTHLLKSLEAQGF $\tt TtAAaAGtGTaGTaGCTGTtATaGGatTGCCgAAtGAtCCctcaGTaCGCATGCAtGAsG$ AaTTtTCaCAtCAtCGACAaTAtCCtaACGGcTTaCTaGGgagtCAtGCGTACGTaCTtC K S V V A V I G L P N D P S V R M H E A CtCTtGGATATGCtCCaGaGGtATGtTGaGGGCaGCaGGtTTCAAaCAtGGaAAtTGGC GaGAaCCTATACGaGGgtCtCCaTACaACtCCCGtCGtCCaAAGTTtGTaCCtTTaACCG LGYAPRGMLRAAGFKHGNWH ATGAtGTaGGTTTtTGGCAaCTtGAcTTCtcttTaCCaGTACCtCCtCGTCccGTttTaC TACTaCAtCCAAAaACCGTtGAaCTgAAGagaaAtGGtCATGGaGGaGCAGGgCAaaAtG D V G F W Q L D F S L P V P P R P V L P Bolif

. .

Figure 20B

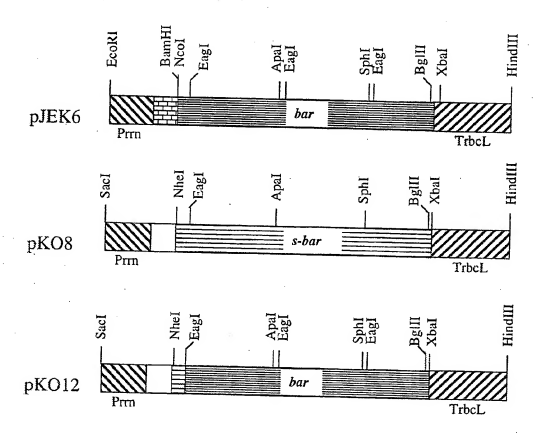


Figure 21

Bacterial Extracts

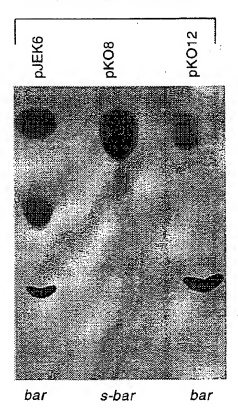
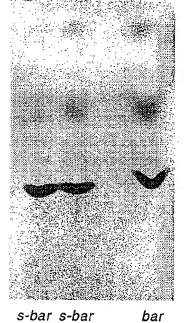


Figure 22A

Plant Extracts



bar

Figure 22B

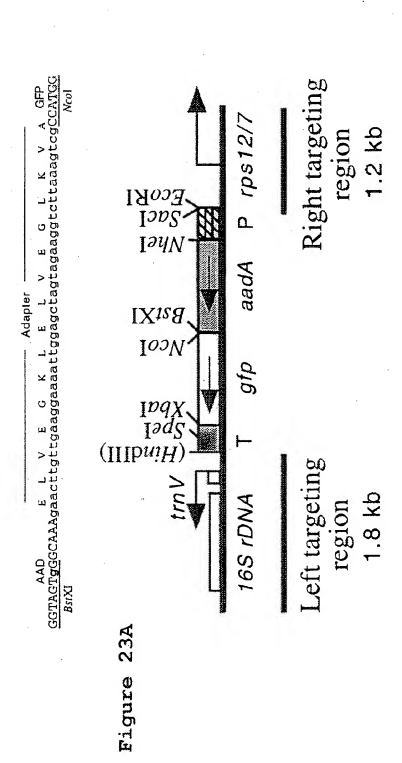


Figure 23E

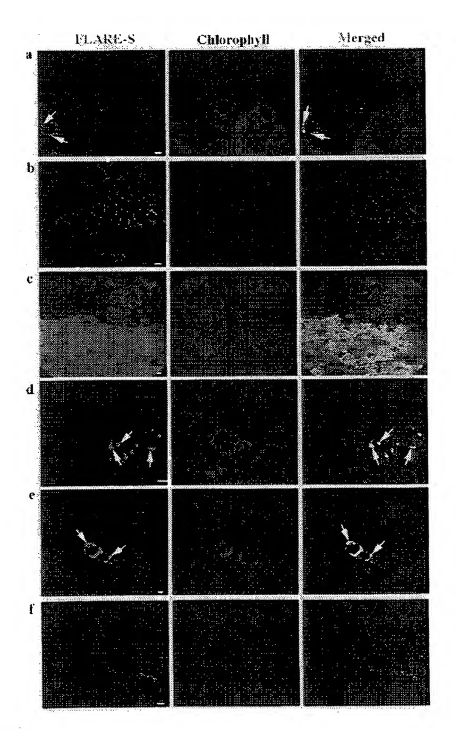


Figure 24

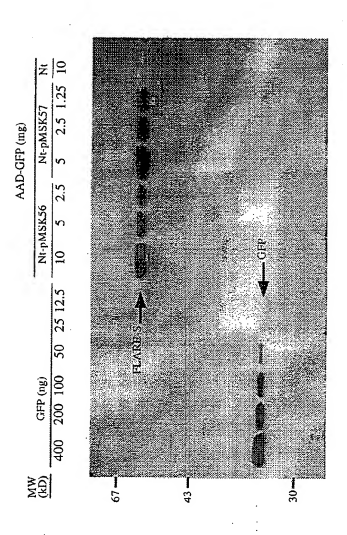


Figure 25

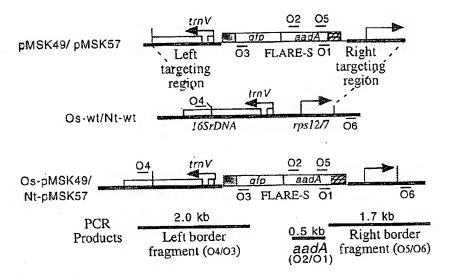


Figure 26A

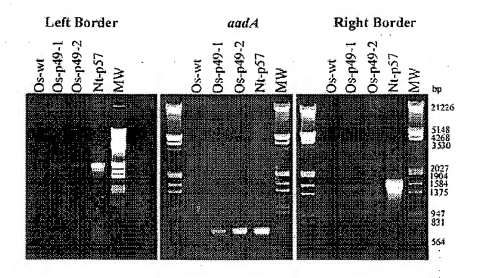


Figure 26B

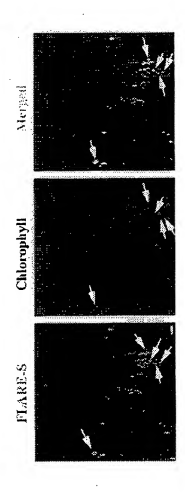


Figure 27

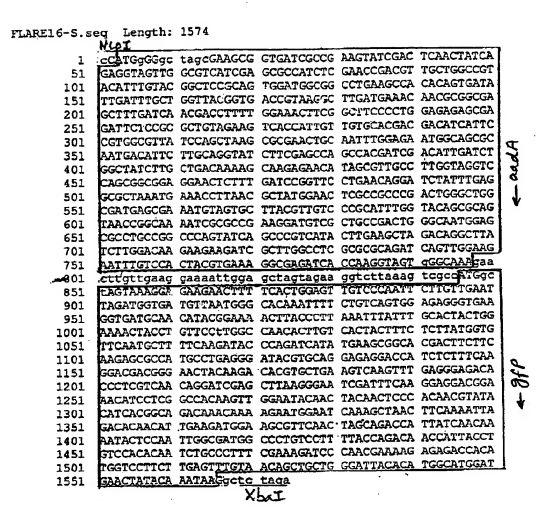


Figure 28

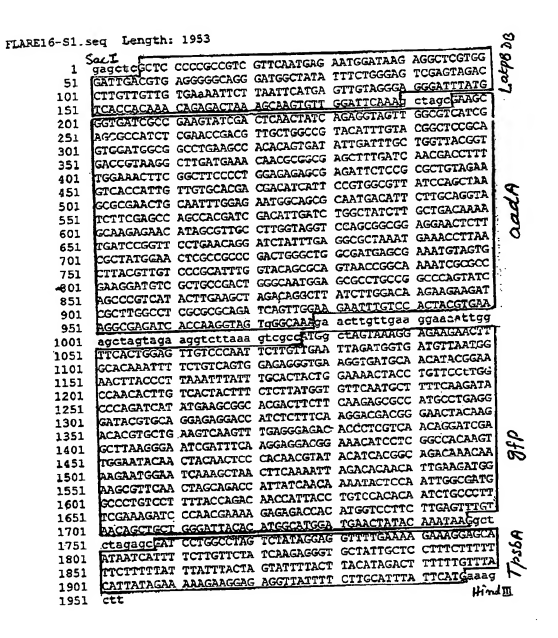


Figure 29

FLARE16-S2.seq Length: 1985

	Sact	
1		8
51	GATTGACGTG AGGGGGCAGG GATGGCTATA TTTCTGGGAG AATTAACCGA	2
101	TCGACGTGCA AGCGGACATT TATTTTAAAT TCGATAATTT TTGCAAAAAC	rbel
151		#
201	TOTGETTOTG GGGTTTCCAC GgctagcGAA GCGGTGATCG CCGAAGTATC	~
251	GACTCAACTA TCAGAGGTAG TTGGCGTCAT CGAGCGCCAT CTCGAACCGA	Í
301	CGTTGCTGGC CGTACATTTG TACGGCTCCG CAGTGGATGG CGGCCTGAAG	İ
351	CCACACAGTG ATATTGATTT GCTGGTTACG GTGACCGTAA GGCTTGATGA	ŀ
401	AACAACGCGG CGAGCTTTGA TCAACGACCT TTTGGAAACT TCGGCTTCCC	Í
451	CTGGAGAGAG CGAGATTCTC CGCGCTGTAG AAGTCACCAT TGTTGTGCAC	ĺ
501	GACGACATCA TTCCGTGGCG TTATCCAGCT AAGCGCGAAC TGCAATTTGG	
551	AGARTGGCAG CGCARTGACA TICTTGCAGG TATCTTCGAG CCAGCCACGA	T
601	TCGACATTGA TCTGGCTATC TTGCTGACAA AAGCAAGAGA ACATAGCGTT	7
651	GCCTTGGTAG GTCCAGCGGC GGAGGAACTC TTTGATCCGG TTCCTGAACA	8
701	GGATCTATTT GAGGCGCTAA ATGAAACCTT AACGCTATGG AACTCGCCGC	ď
751	CCGACTGGGC TGGCGATGAG CGAAATGTAG TGCTTACGTT GTCCCGCATT	_
~ 801	TEGTACAGCE CAGTAACCEE CAAAATCECE CCGAAGGATE TCGCTGCCGA	
851	CTGGGCAATG GAGCGCCTGC CGGCCCHGTA TCAGCCCGTC ATACTTGAAG	
901	CTAGACAGGC TTATC11GGA CAAGAAGAAG ATCGCTTGGC CTCGCGCGCA	
951	GATCAGITGG AAGAATTTGT CCACTACGTG AAAGGCGAGA TCACCAAGGT	
1001	AGTGGGCAAP gaacttgttg aaggaamatt ggagctagta gaaggtctta	
1051	aagtegecAT GgetAGTAAA GGAGAAGAAC TTTTCACTGG AGTTGTCCCA	
1101	ATTCTTGTTG AATTAGATGG TGATGTTAAT GGGCACAAAT TTTCTGTCAG	
1151	TGGAGAGGGT GAAGGTGATG CAACATACGG AAAACTTACC CTTAAATTTA	
1201	TTTGCACTAC TGGAAAACTA CCTGTTCCLT GGCCAACACT TGTCACTACT	
1251	TTCTCTTATG GTGTTCAATG CTTTTCAAGA TACCCAGATC ATATGAAGCG	
1301	GCACGACTTC TTCAAGAGCG CCATGCCTGA GGGATACGTG CAGGAGAGGA	
1351	CCATCTCTTT CAAGGACGAC GGGAACTACA AGACACGTGC TGAAGTCAAG	
1401	TTTGAGGGAG ACACCCTCGT CAACAGGATC GAGCTTAAGG GAATCGATTT	0
1451	CAAGGAGGAC GGAAACATCC TCGGCCACAA GTTGGAATAC AACTACAACT	the state of
1501	CCCACAACGT ATACATCACG GCAGACAAAC AAAAGAATGG AATCAAAGCT	40
1551	AACTTCAAAA TTAGACACAA CATTGAAGAT GGAAGCGTTC AACTAGCAGA	
1601	CCATTATCAA CAAAATACTC CAATTGGCGA TGGCCCTGTC CTTTTACCAG	
1651	ACAACCATTA CCTGTCCACA CAATCTGCCC TTTCGAAAGA TCCCAACGAA	
1701	AAGAGAGACC ACATGGTCCT TCTTGAGTTT GTAACAGCTG CTGGGATTAC	
1751	ACATGGCATG GATGAACTAT ACAAATAAGg ctctagaggs ATCCTGGCCT	Ġ
1801	AGTCTATAGG AGGTTTTGAA AAGAAAGGAG CAATAATCAT TTTCTTGTTC	psba
1851	TATCAAGAGG GIGCTATIGC ICCTITCTIT ITTICTITIT ATTIATITAC	2
1901	TAGTATTTA CTTACATAGA CTTTTTTGTT TACATTATAG AAAAAGAAGG	ド
1951	AGACGTTATT TTCTTGCATT TATTCATGAE agctt	
	Hmd III	

Figure 30

FLARE11-S.seq Length: 1595 C- WAC NCOI CCAtgggggc tagcgaacaa aaactcattt ctgaagaaga cttgcctagc GAAGCGGTGA TCGCCGAAGT ATCGACTCAA CTATCAGAGG TAGTTGGCGT 101 CATCGAGCGC CATCTCGAAC CGACGTTGCT GGCCGTACAT TTGTACGGCT 151 CCGCAGTGGI. TGGCGGCCTG AAGCCACACA GTGATATTGA TTTGCTGGTT ACGGTGACCG TAAGGCTTGA TGAAACAACG CGGCGAGCTT TGATCAACGA 251 CCTTTTGGAA ACTTCGGCTT CCCCTGGAGA GAGCGAGATT CTCCGCGCTG TAGAAGTCAC CATTGTTGTG CACGACGACA TCATTCCGTG GCGTTATCCA 301 GCTAAGCGCG AACTGCAATT TGGAGAATGG CAGCGCAATG ACATTCTTGC 351 AGGTATCTTC GAGCCAGCCA CGATCGACAT TGATCTGGCT ATCTTGCTGA CAAAAGCAAG AGAACATAGC GTTGCCTTGG TAGGTCCAGC GGCGGAGGAA CTCTTTGATC CGGTTCCTGA ACAGGATCTA TTTGAGGCGC TAAATGAAAC 401 451 501 551 CTTAACGCTA TGGAACTCGC CGCCCGACTG GGCTGGCGAT GAGCGAAATG TAGTGCTTAC GTTGTCCCGC ATTTGGTACA GCGCAGTAAC CGGCAAAATC GCGCCGAAGG ATGTCGCTGC CGACTGGGCA ATGGAGCGCC TGCCGGCCCA GTATCAGCCC GTCATACTTG AAGCTAGACA GGCTTATCTT GGACAAGAAG 751 AAGATCGCTT GGCCTCGCGC GCAGATCAGT TGGAAGAATT TGTCCACTAG -801 GTGAAAGGCG AGATCACCAA GGTAGTGGGC AAAbaacttg Cagttgaagg 851 aaaattggag gtcgccATGg ctAGTAAAGG AGAAGAACTT TTCACTGGAG TTGTCCCAAT TCTTGTTGAA TTAGATGGTG ATGTTAATGG GCACAAATTT TCTGTCAGTG GAGAGGGTGA AGGTGATGCA ACATACGGAA AACTTACCCT 1001 TAAATTTATT TGCACTACTG GAAAACTACC TGTTCCCTGG CCAACACTTG TCACIACTIT CTCTTATGGT GTTCAATGCT TTTCAAGATA CCCAGATCAT ATGAAGCGGC ACGACTTCTT CAAGAGCGCC ATGCCTGAGG GATACGTGCA 1051 1101 1151 GGAGAGGACC ATCTCTTCA AGGACGACGG GAACTACAAG ACACGTGCTG 1201 AAGTCAAGTT TGAGGGAGAC ACCCTCGTCA ACAGGATCGA GCTTAAGGGA ATCGATTTCA AGGAGGACGG AAACATCCTC GGCCACAAGT TGGAATACAA 1301 CTACAACTCC CACAACGTAT ACATCACGGC AGACAAACAA AAGAATGGAA 1351 TCAAAGCTAA CTTCAAAATT AGACACAACA: TTGAAGATGG AAGCGTTCAA 1401 CTAGCAGACC ATTATCAACA AAATACTCCA ATTGGCGATG GCCCTGTCCT 1451 TTTACCAGAC AACCATTACC TGTCCACACA ATCTGCCCTT TCGAAAGATC CCAACGAAAA GAGAGACCAC ATGGTCCTTC TTGAGT<u>TTGT AACAGCTGCT</u> 1551 GGGATTACAC ATGGCATGGA TGAACTATAC AAATAAGgct ctaga Xbai

Figure 31

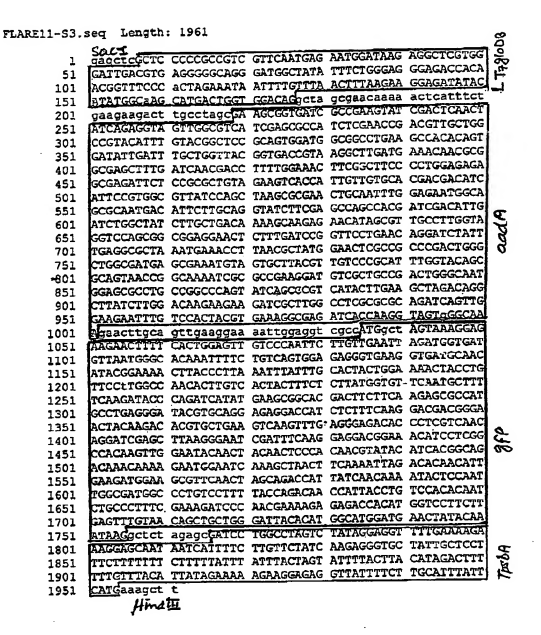


Figure 32

pMSK35.seq Length: 4671 Figure 33A

o 122 488 1 JGGGAACGGAT TCACCGCCGT ATGGCTGACC GGCGATTACT AGCGATTCCT GCTTCATGCA GGCGAGTTGC AGCCTGCAAT CCGAACTGAG GACGGGTTTT 51 TGGAGTTAGC TCACCCTCGC GAGATCGCGA CCCTTTGTCC CGCCCATTGT 101 151 AGCACGTGTG TCGCCCAGGG CATAAGGGGC ATGATGACTT GGCCTCATCC 201 TCTCCTTCCT CCGCCTTAAC ACCGGCGGTC TGTTCAGGGT TCCAAACTCA 251 TAGTGCCAAC TAAACACGAG GGTTGCGCTC GTTGCGAGAC TTAACCCAAC 301 ACCTTACGGC ACGAGCTGAC GACAGCCATG CACCACCTGT GTCCGCGTTC 351 CCGAGGGCAC CCCTCTCTT CAAGAGGATT CGCGGCATGT CAAGCCCTGG TAAGGTTCTT CGCTTTGCAT CGAATTAAAC CACATGCTCC ACCGCTTGTG 401 CGGGCCCCCG TCAATTCCTT TGAGTTTCAT ICTTGCGAAC GTACTCCCCA 451 501 GGCGGGATAC TTAACGCGTT AGCTACAGCA CTGCACGGGT CGAGTCGCAC AGCACCTAGT ATCCATCGTT TACGGCTAGG ACTACTGGGG TCTCTAATCC 551 601 CATTTGCTCC CCTAGCTTTC GTCTCTCAGT GTCAGTGTCG GCCCAGCAGA 651 GTGCTTTCGC CGTTGGTGTT CTTTCCGATC TCAATGCATT TCACCGCTCC 701 ACCGGAAATT CCCTCTGCCC CTACCGTACT CCAGCTTGGT AGTTTCCACC 2 GCCTGTCCAG GGTTGAGCCC TGGGATTTGA CGGCGGACTT GAAAAGCCAC 751 CTACAGACGC TTTACGCCCA ATCATTCCGG ATAACGCTTG CATCCTCTGT CTTACCGCGG CTGCTGGCAC AGAGTTAGCC GATGCTTATT CCTCAGATAC CGTCATTGTT TCTTCTCCGA GAAAAGAAGT TGACGACCCG TGGGCCTTCC 901 ACCTCCACGC GGCATTGCTC CGTCAGGCTT TCGCCCATTG CGGAAAATTC 1001 CCCACTGCTG CCTCCCGTAG GAGTCTGGGC CGTGTCTCAG TCCCAGTGTG 1051 GCCTCACCAA CTAGCTAATC AGACGCGAGC CCCTCCTTGG GCGGATTTCT CCTTITGCTC CTCAGCCTAC GGGGTATTAG CAACCGTTTC CAGTTGTTGT TCCCCTCCCA AGGGCAGGTT CTTACGCGTT ACTCACCCGT TCGCCACTGG 1151 1201 1501 ATCCATTCCC GTTCGATCGT GGCGGGGGGA GTAAGTCAAA ATAGAAAAAA 🗸 1551 CTCACATTGG GTTTAGGGAT AATCAGGCTC GAACTGATGA CTTCCACCAC 1601 GTCAAGGTGA CACTCTACCG CTGAGTTATA TCCCTTCCGC GTCCCCTCGA GRAAGAGAAT TACCGAATCC TAAGGCAAAG GGGCGAGAAA CTCAAGGCCA CCCTTCCTCC GGGCTTTCTT TCCACACTAT TATGGATAGT CAAATAATGG 1651 1701 GAAAAATTGG ATTCAATTGT CAACCGGTCC TATCGAAAAT AGGATTGACT ATGGATTCGA GCCATAGCAC ATGGTTTCAT AAAATCTGTA CGATTTTCCC 1851 GATCTAAATC GAGCAGGTTT CCATGAAGAR gatcgacggt atcgataagc 1901 ttgcatgcct gcaggtCGAA TATAGCTCTT CTTTCTTATT TCAATGATAT 1951 TATTATTTCA AAGATAAGAG ATATTCAAAG ATAAGAGATA AGAAGAAGTC 2001 AAAATTTGAT TTTTTTTTTG GAAAAAAAA ATCAAAAAGA TATAGTAACA 2051 TTAGCAAGAA GAGAAACAAG TTCTATTCA CAATTTAAAC AAATACAAAA 2101 TCAAAATAGA ATACTCAATC ATGAATAAAT GCAAGAAAAT AACCTCTCCT 2151 TCTTTTCTA TAATGTAAAC AAAAAAGTCT ATGTAAGTAA AATACTAGTA 2201 AATAAATAAA AAGAAAAAAA GAAAGGAGCA ATAGCACCCT CTTGATAGAA 2251 CAAGAAATG ATTATTGCTC CITTCTTTTC AAAACCTCCT ATAGACTAGG
2301 CCAGGATOgc tctagcTAGA CATTATTTGC CGACTACCTT GGTGATCTCG 2301 CCAGGATOGE tetageTAGA CATTATTTGE COACIACTIC GEGAGGECIA COTTTCACGT AGTGGACAAA TTCTTCCAAC TGATCTGCGC GCGAGGCCAA TCTTCTCCAA GATAAGCCTG TCTAGCTTCA AGTATGACGG 2451 GCTGATACTG GGCCGGCAGG CGCTCCATTG CCCAGTCGGC AGCGACATCC 2501 TTCGGCGCGA TTTTGCCGGT TACTGCGCTG TACCAAATGC GGGACAACGT 2551 AAGCACTACA TTTCGCTCAT CGCCAGCCCA GTCGGGCGGC GAGTTCCATA 2601 GCGTTAAGGT TTCATTTAGC GCCTCAAATA GATCCTGTTC AGGAACCGGA
2651 TCAAAGAGTT CCTCCGCCGC TGGACCTACC AAGGCAACGC TATGTTCTCT
2701 TGCTTTTGTC AGCAAGATAG CCAGATCAAT GTCGATCGTG GCTGGCTCGA

pMSK35.seq Length: 4671

2751	AGATACCTGC	AAGAATGTCA	TTGCGCTGCC	ATTCTCCAAA	TTGCAGTTCG	î
2801	CGCTTAGCTG	GATAACGCCA	CGGAATGATG	TCGTCGTGCA	CAACAATGGT	
2851		GCGCGGAGAA		TCCAGGGGAA	GCCGAAGTTT	12
2901	CCAAAAGGTC	GTTGATCAAA	GCTCGCCGCG	TTGTTTCATC	AAGCCTTACG	1 2
2951	GTCACCGTAA	CCAGCAAATC	AATATCACTG		GGCCGCCATC	7
3001	CACTGCGGAG	CCGTACAAAT	GTACGGCCAG	CAACGTCGGT	TCGAGATGGC	1
3051		GCCAACTACC		GAGTCGATAC	TTCGGCGATC	•
3101	ACCECTTCCC	TCATGGATCC	CTCCCTACAA	CTGTATCCAa	GCGCTTCgTA	į
3151	TTCGCCCGGA	GTTCGCTCCC	AGAAATATAG	CCATCCCTGC	CCCCTCACGT	İ
3201	CAATCCCACG	AGCCTCTTAT	CCATTCTCAT	TGAACGACGG	CGGGGGAGO	
3251			cctgcagccc			ŧ
3301	AACTGGGGCT	ACATTTCTTT	TCAATTTCCA	TTCAAGAGTT	TCTTLICTGT	1
3351	TTCCACGCCC	TTTTTTGAGA	CCTCGAAACA	TGAAATGGAC	AAATTCCTTC	
3401			AAAAGGATAA			ena
3451	ACTTCATTTC	ATTTATGAAT	TTCATAGTAA	TAGAAATCCA	TETCCTACCE	3
3501	AGACAGAATT	TCGAACTTGC	TATCCTCTTG	CCTAATAGGC	AAAGATTGAC	16
3551	CTCTGTAGAA	AGAATGATTC	ATTCGGATCG	ATATGAGGAC	CCAACTACGT	de
3601	TGCATTGCAG	AATCCATGTT	CCATATTTGA	AGAGGGTTGA	CCTCTGTGCT	3
-26 51			TTCCTGCTGA			8
3701	ACAGAGAAAA	AATGGAGGAC	TGGTGCCGAC	AGTTCATCAC	GGAAGAAAGA	23
3751	ACTCACAGAG	CCGGGATCGC		AATAGTACTA		13
3801	CTAATATATA	GAAATAGATA	TctagctagA	AATAGAAACA	ACTAATATAT	Ø
3851	AGATAATCGA	AATTGAAAAG	AACTGTCTTT	TCTGTATACT	TTCCCCGTTC	82
3901			TGCAATCGAT			2
3951	TCAACACAAC	ATAGGTCATC	GAAAGGATCT			1
4001	GAAAGCCAGT	TAGAAAATGG		Gaagagtgcc		7
4051		CATTAACCCG		ATCCAATTCG		B
4101	TGGGAAGTTT	CGGGAAGAAA	TTGGAATGGA			2
4151		CTCTATTGAT		TACCTAGAGG		*
4201	AGGAAGAGGG.	.AAAAATCGAA	ATGAAATAAA	TAAAGAATAA	AGCAAAAAA	(O
4251			AGCCCAGATT			,9
4301	TCGAAAAGGA	TCCTTCTGAT		TGAGGGGCAA		X
4351	ACCGAGAAAG	ATTTCTTCTT	ATTATAAGAC	•		
4401	TTGGTAAAAG	AACAATCTTC		ATAAATGGAA		
4451	TAGAACATGA		CAATTGGTCT		TCGGGACGGA	
4501	GTGGAAGAAA		TCTCGAACGA		CCCTTCGAAA	
4551	GAATTGAACG			ATCTCATGTA		
4601	GAGGGACAGG		TATCTGTCGA	CTTTTCCACT	ATCAACCCCA	
4651	AAAAACCCAA	CTCTGCCTTA	<u>a</u>			

125878

Figure 33B

pMSK49.seq Length: 5263

Figure 34A

	122488					
1	GGGAACGGAT	TCACCGCCGT	ATGGCTGACC	GGCGATTACT	AGCGATTCCT	
51	GCTTCATGCA	GGCGAGTTGC	AGCCTGCAAT	CCGAACTGAG	GACGGGTTTT	
101	TGGAGTTAGC	TCACCCTCGC	GAGATCGCGA	CCCTTTGTCC	CGCCCATTGT	
151	AGCACGTGTG	TCGCCCAGGG	CATAAGGGGC	ATGATGACTT	GGCCTCATCU	
201	TCTCCTTCCT	CCGGCTTAAC	ACCGGCGGTC	TGTTCAGGGT	TCCAAACTCA	
251	TAGTGGCAAC	TAAACACGAG	GGTTGCGCTC	GTTGCGAGAC	TTAACCCAAC	
301	ACCTTACGGC	ACGAGCTGAC	GACAGCCATG	CACCACCTGT	GTCCGCGTTC	
351	CCGAGGGCAC	CCCTCTCTTT	CAAGAGGATT	CGCGGCATGT	CAAGCCCTGG	d
401	TARGETTETT	CGCTTTGCAT	CGAATTAAAC	CACATGCTCC	ACCGCTTGTG	Ü
451	ceeecccee	TCAATTCCTT	TGAGTTTCAT	TCTTGCGAAC	GTACTCCCCA	3
501	GCCGGGATAC	TTAACGCGTT	AGCTACAGCA	CTGCACGGGT	CGAGTCGCAC	É
551	ACCACCTACT	ATCCATCGTT	TACGGCTAGG	ACTACTGGGG	TCTCTAATCC	à
601	CAMMICCICC	CCTAGCTTTC	GTCTCTCAGT	GTCAGTGTCG	GCCCAGCAGA	0
651	CTCCTTTCCC	CGTTGGTGTT	CTTTCCGATC	TCAATGCATT	TCACCGCTCC	⊘
701	ACCECADATT	CCCTCTGCCC	CTACCGTACT	CCAGCTTGGT	AGTTTCCACC	~
751	CCCTCTCCAG	GGTTGAGCCC	TEGERTTER	CGGCGGACTT	GAAAAGCCAC	Œ.
-601	CENCACACEC	TTTACGCCCA	ATCATTCCGG	ATAACGCTTG	CATCCTCTGT	3
851	CIACAGACGC	CTGCTGGCAC	AGACTTAGCC	GATGCTTATT	CCTCAGATAC	بي
901	CTIACCGCGG	TOTAL GOOD	CANADCAACT	TGACGACCCG	TEGECCTTCC	B
951	ACCECCACC	GGCATTGCTC	CCTCACCCTT	TOGCCCATTG	CGGAAAATTC	Ĺ
1001	ACCI CUACGC	CCTCCCGTAG	CACTCTCCCC	CGTGTCTCAG	TCCCAGTGTG	K
1051	CCCACIGCIG	CTCTCGGACC	ACCUPACTE AT	CATCGCCTTG	GTAAGCTATT	X
	CCCTCACCAA	CTAGCTAATC	VGCTVCLGUL.	CCCTCCTTGG	GCGCATTTCT	dip
1101	CCTUACCAA	CTCAGCCTAC	GEGETATTAG	CAACCGTTTC	CAGTTGTTGT	~~~
1151 1201	DCCCCTCCCD	AGGGCAGSTT	CTTACCCCTT	ACTCACCCGT	TOGCCACTGG	
	AAACACCACT	TCCCGTTCGA	CTTGCATGTG	TTAAGCATGC	CCCCACCGTT	\mathcal{B}_{\cdot}
1251 1301	CATCCTCACC	CAGGATCGAA	CTCTCCATCA	GATTCATAGE	TGCATTACTT	Š
	ATACCTUAGE	TATTCCTAGA	CICICCATOR	TOTGARTIGT	CTTICCTTCC	N.
1351	ALAGCIICCI .	TGTATCCATG	CCANGCAGET	TATTAGCCTG	GAGTTCGCCA	
1401	MAGGAIMACI	AGCCAACCCT	ACCUTATOAC	GTCAATCCCA	CAAGCCTCTT	
1451	CCHGCHGINI	GITCGATCGI	SCCCINICAS.	GTAAGTCAAA	ATAGAAAAA	
1501	MICCALLOCC	GTTTAGGGAT	みなかでおこのでので	CAACTGATGA	CTTCCACCAC	
1551 1601	CTCACALIGG	CACTCTACCG	CTCACTTATA	TOCCTTCCCC	GTCCCCTCGA	
1651	CANAGIGA	TACCGAATCC	TARCCCAARG	GGGCGAGAAA	CTCAAGGCCA	
1701	CCAMCCACC	CGCTTTCTT	TCCACACTAT	TATGGATAGT	CAAATAATGG	
1751	CANADATTICS	ATTCAATTGT	CARCGGTCC	TATCGAAAAT	AGGATTGACT	
1801	ATCCATTCCA	GCCATAGCAC	ATGGTTTCAT	AAAATCTGTA	CGATTTTCCC	
1851	CATCTABATC	GAGCAGGTTT	CCATGAAGAA	CALCGACGGE	atcgataagc	
1901	+++CATCAAT	AAATGCAAGA	AAANAACOIC	TECTTCTTTT	TCTATAATGT	~
1951	DADCABABA	GTCTATGTAA	GTAAAATACT	AGTAAATAAA	TAAAAAGAAA	-Š
2001	DANGARAGE	AGCAATAGCA	CCCTCTTGAT	AGAACAAGAA	AATGATTATT	12
2051	المسادر مساسات المساح	ጥጥጥሮች ሽ ሽ ሽ ሽ ር ር ር	TOTATACAC	TAGGCCAGGA	TCoctctaga	•
2101	CO TOTA TOTAL	татасттсат	CCATGCCATG	TGTAATCCCA	GCAGCTGTTA	
2151	CARACTCARG	AAGGACCATG	TEGTCTCTCT	TTTCGTTGGG	ATCTTTCGAA	
2201	ACCCCAGATT	GTGTGGACAG	GTAATGGTTG	TCTGGTAAAA	GGACAGGGCC	
2251	PACCOCAPATA	CCACTATTTT	GTTGATAATG	GTCTGCTAGT	TGAACGCTTC	
2301	CATCTTCAAT	CTTCTCTCTCTA	ATTTTGAAGT	TAGCTTTGAT	TCCATTCTTT	ŀ
2351	reminderence	CCCTCATCTA	TACGTTGTGG	GAGTTGTAGT	TGTATTCCAA	
2401	רדופוכום הדופוכוכום	ACCATCTTTC	CGTCCTCCTT	GAAATCGATT	CCCTTAAGCT	
2451	CATCOTOTT	CACCACCCTC	TCTCCCTCAA	ACTICACTIC	AGCACGTGTC	6
2501	PROTECTION	CGTCGTCCTT	CADAGAGATG	GTCCTCTCCT	GCACGTATCC	38
2551	CTCA CCCATC	GCGCTCTTGA	DEDDETCETE	CCGCTTCATA	TGATCTGGGT	l '
2601	ATCTTCAAAA	CCATTCAACA	CCATARGE	AAGTAGTGAC	AAGTGTTGGC	
2651	CAPCCAPCAC	ChyChumutcc	DETACTORA	ATAAATTTAA	GGGTAAGTTT	1
	haden action	GLEGITIO	CACCCTCTCC	ACTGACAGAA	AATTTGTGCC	ł
2701	"CCGINIGII	and touch I	-	· · · · · · · · · · · · · · · · · · ·		-

Figure 34B

pMSK49.seq Length: 5263

		_			TCCACTGAAA	
2751	CATTAACATC	ACCATCTAAT	TCAACAAGAA	1114446	cttcaactgc	
2801						
2851	aaqttcTTTG	CCCACTACCT	TGGTGATCTC	GCCTTTCACG	TTCTTGTCCA	
2901						
2951						
3001		COCCA COCCC	CACCCACACC	CITLEGUCGUG	WTTYV	
3051			CYPECACAACI	TAMECALCARC	*** * * * * * * * * * * * * * * * * * *	
3101		* WEACCCCCC	CAL VILLAL A. V. V.	A(-('PT TUNDA	~	
3151			CACCAACCE	ATCAMMOMUL	TOOTOGGGGGG	. ·
3201		~~ * ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	ر) البراياتياتي تدلايات	TIGUTITIES	C470-C4 N	7
3251	C3 MC3 3	かんせんにかかしたか		AAGATACCTG	Of Milorania and al	V
3301	L-managemen		Villa La Villa Carlon C		COURT STREET	g
3351	ACGGAATGAT	GTCGTCGTGC	ACAACAATGG	TGACTTCTAC	ACCOCCOCACA ACCA A	U
3401		ベルベベス かかかたる	ACCCCAACIIII	TOURNAMENT	COLLONIA	
3451		بالركز والمنطوطات باستنت	CAAGCCTTTAC	GGILHUUGIA	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
3501	I		ACCOUNT OF THE	CCALIGORA	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
3551	1		田中へて れたれが(こ)こ	CCTCGATGA		
3601		あっち ごでこころ 中の		CACCIGUITA	Cred acada	28
-2651			*****************		The passes and a	
3701						20
3751						
3801						}\ [
3851			CCCATCITAL	COTTACONS.		7
3901		Albuthath W. W. Albuthath	CATTCAMGAG	TITOTIMA	~	
3951			CATGAAATGG	ACAMATICCE	TOTOTTOO	1 3
4001	ACACATACAA	GAAAAAGGAT	AATGGTAGCC	CTCCCATTAA	CONCRETE	13
4051	t	<u> ል ውውው ምክ ምክ ርግ</u>	カカザカにカカムがに	LAIGILLIAN	CONTRACTOR	3
4101			- ובייים בייים בייים	GITHOWAY:P	TICAT OF A **	
4151				DETERMENT	GTIGMITE	1 0
4201	L				~***	1
4251	1		(CA/2('Y' T T T ')')	TUTULIUGU	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	100
4301	AAAATGGAGG	ACTGGTGCCG	ACAGTTCATC	ACGGAAGAAA	מיים מיים מיים מיים מיים	13
4351	AGCCGGGATC	GCTAACTAAT	AGAATAGTAC	TACTAACTAA	DTD CATABTO	get!
4401			#XXXXTA(_6666	LAALINALAL	17770-0	\$
4451	KARAMME E E E	יוד) וביות ביות או מיבוע	THEFT		PATTE POPE	1 .
4501	CCGCGGGTCT	TATGCAATCG	ATCGGATCAT	ATAGATATCC	ACCARACCE] *
4551	ACATAGGTCA	TCGAAAGGAT	CTCGGACGAC	TCACCAAAGC	TGGATAAGCT	
4601	GTTAGAAAAT	GGATTCCTAT	TIGAAGAGIG	CCTAACCGCA	CTTGGGAAGI	12
4651	CACATTAACC	CGTCAATTTT	GGATCCAATT	CGGGATTTTT	CACCALAAGO	1.05
4701	TTCGGGAAGA	AATTGGAATG	GAATAATATA	GATTCATACA	DELGGAAGAG	0
4751	TTCTCTATTG	ATGCAAACGC	TGTACCTAGA	GGATAGGGAT	AAAAATAAGI	
4801	GGAAAAATCG	AAATGAAATA	AATAAAGAAT	AAAGCAAAAA	ACTCCADAD(8
4851	CGAAGATAGA	AGAGCCCAGA	TTCCAAATGA	AGAAATGGAA	ACTOMISMA ATTOCCOCACA	11.2
4901		- カボケケマンス みんこう	AATCAGGGC	ARUGUCHA I C	MINCOMICA	·170
4951		ם על על על על על על אייני ווייים אייניים אייניים אייניים אייניים אייניים אייניים אייניים אייניים אייניים אייני	LCGTGATITE	ATCCCCAMANA	GILIGGIANG	-1
5001	I CARCARDO	ፈ <i>ጀ</i> ሞሞንጋተንጥ ነ	TCATAAATGG	AAAGTGTTC	WITHGUAGE.	- 1
5051	1	へってって カ のでくくご	بانت لا باربانت لا بابتلس م	CTTCGGGACG	- GWGIGGWAG	3 .
5101	hacecenae	・ みんかいかんごみなく	: GAGGAAAAGG	ATCCCTTCG	7 WHOWNTION	3- J
5151	1-03 003 0000	ተከተመ አርርፕር ጀ	ANATICICATO	TACGALICAC	. Tuckocmie	· - [
5201	GGAAGGGTGA	CTTATCTGTC	GACTITICCA	CTATCAACCO	CAMADACC	
5251	AACTCTGCCT	TAC				

Gene	Product	Plasmid
aadA16gfp	FLARE16-S	pMSK51 (BS)
aadA16gfp-S1	FLARE16-S1	pM\$K56 (Nt-pRV111B)
aadAl6gfp-S2	FLARE16-S2	pMSK57 (Nt-pRV111B)
aadA11gfp-S3	FLARE11-S3	pMSK49 (Os-pMSK49)

Figure 35

International application No. PCT/US99/17806

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A01H 4/00, 5/00; C12N 15/29, 15/31, 15/65, 15/67, 15/82, 15/84 US CL :Please See Extra Sheet.				
According to	cording to International Patent Classification (IPC) or to both national classification and IPC			
	DS SEARCHED			
	cumentation searched (classification system followed	by classification symbols)		
	Please See Extra Sheet.			
Documentati	on searched other than minimum documentation to the	extent that such documents are included	in the fields searched	
Electronic d	ata base consulted during the international search (na	me of data base and, where practicable,	search terms used)	
EMBL, G	· ·	:		
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.	
Y	WO 97/06250 A1 (RUTGERS UNIVERSES 1, 3-6, 12-14, 17-18 and Figures		1-6 and 15-17	
Y	NIELSEN et al. Characterization of the Pea Chloroplast DNA OriA Region. Plasmid. 1993, Vol. 30, pages 197-211, see page 197.			
Y	SEKIYA et al. Sequence of the gene surrounding region in a ribosomal RNA Nucleic Acids Research. 1979, Vol. 575.	A operon of Escherichia coli.	1-6	
	ner documents are listed in the continuation of Box C	. See patent family annex.		
		"I" later document published after the int	ernational filing date or priority	
·A· do	ecial categories of cited documents: cument defining the general state of the art which is not considered	date and not in conflict with the app the principle or theory underlying th	lication but cited to understand	
to	be of particular relevance rlier document published on or after the international filing date	"X" document of particular relevance; the	se claimed invention cannot be	
do	cument which may throw doubts on priority claim(s) or which is and to establish the publication date of another citation or other	when the document is taken alone		
.O. qo	orial reason (as specified) cument referring to an oral disclosure, use, exhibition or other sams	"Y" document of particular relevance, it considered to involve an inventive combined with one or more other such being obvious to a person skilled in	to step when the document is the documents, such combination	
·P· do	cument published prior to the international filing date but later than s priority date claimed	*&* document member of the same pater	at family	
	actual completion of the international search	Date of mailing of the international se	arch report	
07 NOVE	MBER 1999	30 NOV 1999		
Box PCT	Washington, D.C. 20231			

International application No. PCT/US99/17806

	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category*	Chauon of document, with indication, where appropriate, or the following personal	
Y	SPRENGART et al. The downstream box: an efficient and independent translation intitiation signal in <i>Escherichia coli</i> . The EMBO Journal. 1996, Vol. 15, No. 3, pages 665-674, see page 665.	1-6
Y	VASIL et al. Herbicide resistant fertile transgenic wheat plants obtained by microprojectile bombardment of regenerable embryogenic callus. Bio/Technology. June 1992, Vol. 10, No. 6, pages 667-674, see page 667.	7-8 and 18-26
Y	PANG et al. An Improved Green Fluorescent Protein Gene as a Vital Marker in Plants. Plant Physiology. 1996, Vol. 112, pages 893-900, see page 893.	9-14
Y	HIRATSUKA et al. The complete sequence of the rice (Oryza sativa) chloroplast genome: Intermolecular recombination between distinct tRNA genes accounts for a major plastid DNA inversion during the evolution of cereals. Molecular and General Genetics. 1989, Vol. 217, pages 185-194, see page 185.	18-26
Y	ZHANG et al. Efficient regeneration of transgenic plants from rice protoplasts and correctly regulated expression of the foreign gene in plants. Theoretical and Applied Genetics. 1988, Vol. 76, pages 835-840, see page 835.	18-23
Y	THOMPSON et al. Protoplast culture of rice (Oryza sativa L.) using media solidified with agarose. Plant Science. 1986, Vol. 47, No. 2, pages 123-133, see page 123.	24-26
Y	MAKRIDES, S.C. Strategies for Achieving High-Level Expression of Genes in <i>Escherichia coli</i> . Microbiological Reviews. September 1996, Vol. 60, No. 3, pages 512-538, see page 524.	27-28
Y	MURRAY et al. Codon usage in plant genes. Nucleic Acids Research. 1989, Vol. 17, No. 2, pages 477-498, see pages 477 and 491.	27-28
	:	
	:	

International application No. PCT/US99/17806

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all scarchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

International application No. PCT/US99/17806

A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

435/69.7, 320.1, 424, 430, 430.1, 431, 468, 469, 470; 536/23,6, 23.7, 24.1; 800/278, 287, 288, 292, 293, 294, 300, 300.1, 320, 320.1, 320.2, 320.3

B. FIELDS SEARCHED
Minimum documentation searched
Classification System: U.S.

435/69.7, 320.1, 424, 430, 430.1, 431, 468, 469, 470; 536/23,6, 23.7, 24.1; 800/278, 287, 288, 292, 293, 294, 300, 300.1, 320, 320.1, 320.2, 320.3

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s) 1-26, drawn to a DNA construct comprising a promoter element, a leader, and a downstream box element, and a method for its use for enhanced plastid expression of a heterologous gene.

Group II, claim(s) 27-28, drawn to a method for modifying codon usage in structural genes.

The inventions listed as Groups I and II do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

The inventions of each of Groups I, drawn to a first product and method of use, and Group II, drawn to a second method, are taught or suggested by the prior art, as outlined in the accompanying Search Report, and so each do not constitute an advance over the prior art which would constitute a special technical feature. Furthermore, the various leader sequences and downstream box elements of Group I, and the methods for plastid transformation of Group I, are not required by Group II; while the methods of structural gene modification of Group II are not required by Group I. Accordingly, the inventions do not relate to a single inventive concept which would constitute a single special technical feature.